



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:)	Examiner: Saoud, Christine J.
)	
Avi ASHKENAZI, <i>et al.</i>)	Art Unit: 1647
)	
Application Serial No. 09/905,348)	Confirmation No: 3826
)	
Filed: July 13, 2001)	Attorney's Docket No. GNE-1618 P2C18
)	
For: SECRETED AND TRANSMEMBRANE)	Customer No. 77845
POLYPEPTIDES AND NUCLEIC)	
ACIDS ENCODING THE SAME)	

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ON APPEAL TO THE BOARD OF PATENT APPEALS AND INTERFERENCES
APPELLANTS' BRIEF

MAIL STOP APPEAL BRIEF - PATENTS

Commissioner for Patents
P.O. Box 1450
Alexandria, Virginia 32613-1450

Dear Sir:

This Appeal Brief, filed in connection with the above captioned patent application, is responsive to the Final Office Actions mailed on December 27, 2007. A Notice of Appeal was filed on May 27, 2008. The following constitutes the Appellants' Brief on Appeal, submitted with a request for a two month extension of time with the requisite fee. As the two month extension deadline for response fell on a Saturday, this brief is timely filed on the following business day September 29, 2008.

Appellants hereby appeal to the Board of Patent Appeals and Interferences from the final rejection in this case.

I. REAL PARTY IN INTEREST

The real party in interest is Genentech, Inc., South San Francisco, California, by an assignment of the parent application, U.S. Patent Application Serial No. 09/665,350 recorded July 9, 2001, at Reel 011964 and Frame 0181. The present application is a continuation of U.S. Patent Application Serial No. 09/665,350.

II. RELATED APPEALS AND INTERFERENCES

The claims pending in the current application are directed to a polypeptide referred to herein as "PRO232." Although there exist several applications directed to the "gene amplification" utility under Appeal, there are no applications related to PRO232 nucleic acids or antibodies.

III. STATUS OF CLAIMS

Claims 44-46 and 49-51 are in this application.

Claims 1-43 and 47-48 have been canceled.

Claims 44-46 and 49-51 stand rejected and Appellants appeal the rejection of these claims.

A copy of the rejected claims in the present Appeal is provided in the Claims Appendix.

IV. STATUS OF AMENDMENTS

A summary of the prosecution history for this case is as follows:

Previously, in response to a Final Office Action mailed on October 21, 2005, a Notice of Appeal was filed on February 21, 2006 and an Appeal Brief was filed on April 17, 2006. An RCE Response with additional references and affidavits supporting Appellants' arguments was filed on July 27, 2006. A Final Office Action was mailed on October 17, 2006, and a Notice of Appeal was filed on March 19, 2007. An Appeal Brief was filed on September 24, 2007.

Upon further consideration, the Examiner withdrew the finality of the previous Office Action solely to clarify the issues for appeal as outlined in a Final Office Action mailed December 27, 2007. A Notice of Appeal was filed on May 27, 2008.

No claim amendments have been submitted after the last final rejection of October 17, 2006. A copy of the rejected claims in the present Appeal is provided in the Claims Appendix.

V. SUMMARY OF CLAIMED SUBJECT MATTER

The invention claimed in the present application is related to an isolated polypeptide comprising the amino acid sequence of the polypeptide of SEQ ID NO:18, referred to in the present application as “PRO232”. The PRO232 gene was shown for the first time in the present application to be significantly amplified in human lung and colon cancers as compared to normal, non-cancerous human tissue controls (Example 92). This feature is specifically recited in Claims 44(a) and 45, and carried by all claims dependent from Claim 44. In addition, the invention also claims the amino acid sequence of the polypeptide of SEQ ID NO:18, lacking its associated signal-peptide (Claims 44(b) and 46); or the amino acid sequence of the polypeptide encoded by the full-length coding sequence of the cDNA deposited under ATCC accession number 209250 (Claims 44(c) and 49). The invention is further directed to a chimeric polypeptide comprising one of the above polypeptides fused to a heterologous polypeptide (Claim 50), and to a chimeric polypeptide wherein the heterologous polypeptide is an epitope tag or an Fc region of an immunoglobulin (Claim 51). The preparation of chimeric PRO polypeptides (Claims 50 and 51), including those wherein the heterologous polypeptide is an epitope tag or an Fc region of an immunoglobulin, is set forth in the specification at page 74, line 23 to page 75, line 5. Examples 53-56, pages 192-199, describe the expression of PRO polypeptides in various host cells, including *E. coli*, mammalian cells, yeast and Baculovirus-infected insect cells.

The amino acid sequence of the “PRO232” polypeptide and the nucleic acid sequence encoding this polypeptide (referred to in the present application as “DNA34435-1140”) are shown in the present specification as SEQ ID NOs: 18 and 17, respectively, and in Figures 9 and 8, described on page 59, lines 4-7. The full-length PRO232 polypeptide having the amino acid sequence of SEQ ID NO:18 is described in the specification at, for example, on page 4, pages 3-4 and page 100, page 131, lines 9 to 16 and the isolation of cDNA clones encoding PRO232 of SEQ ID NO:18 is described in Example 4, pages 149-150 of the specification. The specification discloses that the PRO232 polypeptide possess significant sequence homology to cell surface stem cell antigen (35% sequence identity with a stem cell surface antigen from *Gallus gallus*) and may play a role in cell proliferation and/or differentiation. (see for example, page 4 and Example 4, lines 14-15).

Finally, Example 92, in the specification at page 222, line 26, to page 235, line 3, sets forth a 'Gene Amplification assay' which shows that the PRO232 gene is amplified in the genome of certain human lung cancers (see Table 9, pages 230-234). The profiles of various primary lung and colon tumors used for screening the PRO polypeptide compounds of the invention in the gene amplification assay are summarized on Table 8, page 227 of the specification.

VI. GROUND OF REJECTION TO BE REVIEWED ON APPEAL

1. Whether instant Claims 44-46 and 49-51 satisfy the utility/enablement requirement under 35 U.S.C. §§101/112, first paragraph.
2. Whether Claims 44-46 and 49-51 are entitled to the priority date of U.S. Provisional Patent Application Serial No. 60/059121, filed September 17, 1997.
3. Whether Claims 44-46 and 49-51 are anticipated under 35 U.S.C. §102(b) by Rosenthal *et al.*, DE19818619-A1 (October 1999).

VII. ARGUMENTS

Summary of the Arguments:

Issue 1: Utility/ Enablement

Appellants rely upon the gene amplification data of the PRO232 gene for patentable utility of the PRO232 polypeptides. This data is clearly disclosed in the instant specification in Example 92 which discloses that the gene encoding PRO232 showed significant amplification, ranging from 2.056-fold to 5.28-fold, in five lung tumors or 2.00-fold to 5.32-fold in seven colon tumors. Example 92 in the instant specification discloses that, "(a)mplification is associated with overexpression of the gene product, indicating that the polypeptides are useful targets for therapeutic intervention in certain cancers such as colon, lung, breast and other cancers and diagnostic determination of the presence of those cancers" (emphasis added).

Appellants have submitted ample evidence to show that, in general, if a gene is amplified in cancer, it is more likely than not that the encoded protein will be expressed at an elevated level. First, Appellants have submitted, in their Response filed July 25, 2005, a Declaration by Dr. Audrey Goddard, which explains that a gene identified as being amplified at least 2-fold by the disclosed gene amplification assay in a tumor sample relative to a normal sample is useful as

a marker for the diagnosis of cancer, and for monitoring cancer development and/or for measuring the efficacy of cancer therapy. Therefore, such a gene is useful as a marker for the diagnosis of lung and colon cancers, and for monitoring cancer development and/or for measuring the efficacy of cancer therapy. Second, Appellants have submitted, in their Responses filed August 9, 2004 and July 27, 2006, ample evidence to show that, in general, if a gene is amplified in cancer, it is more likely than not that the encoded protein will be expressed at an elevated level. For instance, the articles by Orntoft *et al.*, Hyman *et al.*, and Pollack *et al.* collectively teach that in general, gene amplification increases mRNA expression. Third, Appellants have submitted over a hundred references, along with Declarations of Dr. Paul Polakis, which collectively teach that, in general, there is a correlation between mRNA levels and polypeptide levels.

Taken together, although there are some examples in the scientific art that do not fit within the central dogma of molecular biology that there is generally a positive correlation between DNA, mRNA, and polypeptide levels, in general, in the majority of amplified genes, the art overwhelmingly shows that gene amplification influences gene expression at the mRNA and protein levels. Therefore, one of skill in the art would reasonably expect in this instance, based on the amplification data for the PRO232 gene, that the PRO232 polypeptide is concomitantly overexpressed and has utility in the diagnosis of lung and colon cancers.

Appellants further submit that, as evidenced by the Ashkenazi Declaration and the teachings of Hanna and Mornin (both made of record in Appellants' Responses filed December 10, 2003 and August 9, 2004), simultaneous testing of gene amplification and gene product over-expression enables more accurate tumor classification, even if the gene-product, the protein, is not over-expressed. This leads to better determination of a suitable therapy for the tumor, as demonstrated by a real-world example of the breast cancer marker HER-2/neu. Therefore, as a general rule, one skilled in the art would find it more likely than not that PRO232 polypeptides are useful as a diagnostic tools for detecting lung and colon tumors.

As the Examiner no longer questions whether mRNA levels are not predictive of polypeptide levels, the evidence presented by Appellants support that gene amplification correlates with the increased protein expression. Based on the 2.00-fold to 5.32-fold amplification in colon and lung primary tumors, one of ordinary skill would find it credible that

the claimed PRO232 polypeptides would have utility as markers for the diagnosis of lung and colon tumors, and for monitoring cancer development and/or for measuring the efficacy of cancer therapy.

Accordingly, Appellants submit that when the proper legal standard is applied, one should reach the conclusion that the present application discloses at least one patentable utility for the claimed PRO232 polypeptides. Further, one of ordinary skill in the art would understand how to make and use the recited polypeptides for the diagnosis of lung and colon cancers without any undue experimentation.

Accordingly, this enablement rejection under 35 U.S.C. §§101 and 112, first paragraph should be withdrawn.

Issue 2: Priority

The instant application has not been granted the earlier priority date of U.S. Provisional Patent Application Serial No. 60/059121, filed September 17, 1997 on the grounds that the 60/059121 application fails to provide a utility and lacks an enabling disclosure for the claimed invention under 35 U.S.C. §§101/112, first paragraph.

Appellants submit that, for the same reasons discussed above under Issue 1, U.S. Provisional Patent Application Serial No. 60/059121 also satisfies the utility requirements. Therefore, Appellants should be entitled to the priority date of **September 17, 1997**.

Issue 3: Anticipation by Rosenthal *et al.*

As discussed above under Issue 2, the present application should be entitled to the earlier filing date of **September 17, 1997** and therefore, Rosenthal *et al.*, DE19818619-A1, dated October 1999, is not prior art. Thus the instant claims are not anticipated by Rosenthal *et al.*

These arguments are all discussed in greater detail below, under their appropriate headings.

Response to Rejections

ISSUE 1: The Instant Claims 44-46 and 49-51 Satisfy the Utility Requirement under 35 U.S.C. §101/§112, First Paragraph, based on the results of the gene amplification assay

The sole basis for the Examiner's rejection of Claims 44-46 and 49-51 under this section is that the data presented in the instant Application, allegedly, does not satisfy the requirements of 35 U.S.C. §§101/112, first paragraph. Appellants strongly disagree for the reasons discussed below.

A. The Legal Standard For Utility Under 35 U.S.C. §101

According to 35 U.S.C. §101:

Whoever invents or discovers any new and *useful* process, machine, manufacture, or composition of matter, or any new and *useful* improvement thereof, may obtain a patent therefor, subject to the conditions and requirements of this title.
(Emphasis added).

In interpreting the utility requirement, in *Brenner v. Manson*,¹ the Supreme Court held that the *quid pro quo* contemplated by the U.S. Constitution between the public interest and the interest of the inventors required that a patent Applicant disclose a "substantial utility" for his or her invention, *i.e.*, a utility "where specific benefit exists in currently available form."² The Court concluded that "a patent is not a hunting license. It is not a reward for the search, but compensation for its successful conclusion. A patent system must be related to the world of commerce rather than the realm of philosophy."³

Later, in *Nelson v. Bowler*,⁴ the C.C.P.A. acknowledged that tests evidencing pharmacological activity of a compound may establish practical utility, even though they may not establish a specific therapeutic use. The Court held that "since it is crucial to provide researchers with an incentive to disclose pharmaceutical activities in as many compounds as

¹ *Brenner v. Manson*, 383 U.S. 519, 148 U.S.P.Q. (BNA) 689 (1966).

² *Id.* at 534, 148 U.S.P.Q. (BNA) at 695.

³ *Id.* at 536, 148 U.S.P.Q. (BNA) at 696.

⁴ *Nelson v. Bowler*, 626 F.2d 853, 206 U.S.P.Q. (BNA) 881 (C.C.P.A. 1980).

possible, we conclude adequate proof of any such activity constitutes a showing of practical utility."⁵

In *Cross v. Iizuka*,⁶ the C.A.F.C. reaffirmed *Nelson*, and added that *in vitro* results might be sufficient to support practical utility, explaining that "*in vitro* testing, in general, is relatively less complex, less time consuming, and less expensive than *in vivo* testing. Moreover, *in vitro* results with the particular pharmacological activity are generally predictive of *in vivo* test results, *i.e.*, there is a reasonable correlation there between."⁷ The Court perceived, "No insurmountable difficulty" in finding that, under appropriate circumstances, "*in vitro* testing, may establish a practical utility."⁸

The case law has also clearly established that Appellants' statements of utility are usually sufficient, unless such statement of utility is unbelievable on its face.⁹ The PTO has the initial burden to prove that Appellants' claims of usefulness are not believable on their face.¹⁰ In general, an Appellant's assertion of utility creates a presumption of utility that will be sufficient to satisfy the utility requirement of 35 U.S.C. §101, "unless there is a reason for one skilled in the art to question the objective truth of the statement of utility or its scope."^{11, 12}

⁵ *Id.* at 856, 206 U.S.P.Q. (BNA) at 883.

⁶ *Cross v. Iizuka*, 753 F.2d 1047, 224 U.S.P.Q. (BNA) 739 (Fed. Cir. 1985).

⁷ *Id.* at 1050, 224 U.S.P.Q. (BNA) at 747.

⁸ *Id.*

⁹ *In re Gazave*, 379 F.2d 973, 154 U.S.P.Q. (BNA) 92 (C.C.P.A. 1967).

¹⁰ *Ibid.*

¹¹ *In re Langer*, 503 F.2d 1380, 1391, 183 U.S.P.Q. (BNA) 288, 297 (C.C.P.A. 1974).

¹² See also *In re Jolles*, 628 F.2d 1322, 206 USPQ 885 (C.C.P.A. 1980); *In re Irons*, 340 F.2d 974, 144 USPQ 351 (1965); *In re Sichert*, 566 F.2d 1154, 1159, 196 USPQ 209, 212-13 (C.C.P.A. 1977).

Compliance with 35 U.S.C. §101 is a question of fact.¹³ The evidentiary standard to be used throughout *ex parte* examination in setting forth a rejection is a preponderance of the totality of the evidence under consideration.¹⁴ Thus, to overcome the presumption of truth that an assertion of utility by the Appellant enjoys, the Examiner must establish that it is more likely than not that one of ordinary skill in the art would doubt the truth of the statement of utility. Only after the Examiner made a proper *prima facie* showing of lack of utility, does the burden of rebuttal shift to the Appellant. The issue will then be decided on the totality of evidence.

The well established case law is clearly reflected in the Utility Examination Guidelines (“Utility Guidelines”),¹⁵ which acknowledge that an invention complies with the utility requirement of 35 U.S.C. §101, if it has at least one asserted “specific, substantial, and credible utility” or a “well-established utility.” Under the Utility Guidelines, a utility is “specific” when it is particular to the subject matter claimed. For example, it is generally not enough to state that a nucleic acid is useful as a diagnostic without also identifying the conditions that are to be diagnosed.

In explaining the “substantial utility” standard, M.P.E.P. §2107.01 cautions, however, that Office personnel must be careful not to interpret the phrase “immediate benefit to the public” or similar formulations used in certain court decisions to mean that products or services based on the claimed invention must be “currently available” to the public in order to satisfy the utility requirement. “Rather, any reasonable use that an applicant has identified for the invention that can be viewed as providing a public benefit should be accepted as sufficient, at least with regard to defining a ‘substantial’ ‘utility.’”¹⁶ Indeed, the Guidelines for Examination of Applications for Compliance With the Utility Requirement,¹⁷ gives the following instruction to

¹³ *Raytheon v. Roper*, 724 F.2d 951, 956, 220 U.S.P.Q. (BNA) 592, 596 (Fed. Cir. 1983) *cert. denied*, 469 US 835 (1984).

¹⁴ *In re Oetiker*, 977 F.2d 1443, 1445, 24 U.S.P.Q.2d (BNA) 1443, 1444 (Fed. Cir. 1992).

¹⁵ 66 Fed. Reg. 1092 (2001).

¹⁶ M.P.E.P. §2107.01.

¹⁷ M.P.E.P. §2107 II(B)(1).

patent examiners: "If the Applicant has asserted that the claimed invention is useful for any particular practical purpose . . . and the assertion would be considered credible by a person of ordinary skill in the art, do not impose a rejection based on lack of utility."

B. Proper Application of the Legal Standard

Appellants submit that the evidentiary standard to be used throughout *ex parte* examination of a patent application is a preponderance of the totality of the evidence under consideration. Thus, to overcome the presumption of truth that an assertion of utility by the Applicant enjoys, the Examiner must establish that it is more likely than not that one of ordinary skill in the art would doubt the truth of the statement of utility. Only after the Examiner has made a proper *prima facie* showing of lack of utility, does the burden of rebuttal shift to the Applicant.

Appellants respectfully submit that the data presented in Example 92 starting on page 222 of the specification and the cumulative evidence of record support a "specific, substantial and credible" asserted utility for the presently claimed invention.

Patentable utility for the PRO232 polypeptides is based upon the gene amplification data for the gene encoding the PRO232 polypeptide. Example 92 describes the results obtained using a very well-known and routinely employed polymerase chain reaction (PCR)-based assay, the TaqManTM PCR assay, also referred to herein as the gene amplification assay. This assay allows one to quantitatively measure the level of gene amplification in a given sample, say, a tumor extract, or a cell line. It was well known in the art at the time the invention was made that gene amplification is an essential mechanism for oncogene activation. Appellants isolated genomic DNA from a variety of primary cancers and cancer cell lines that are listed in Table 9 (pages 222 onwards of the specification), including primary lung and colon cancers of the type and stage indicated in Table 8 (page 227). The tumor samples were tested in triplicates with TaqmanTM primers and with internal controls, beta-actin and GADPH in order to quantitatively compare DNA levels between samples (page 229). As a negative control, DNA was isolated from the cells of ten normal healthy individuals, which was pooled and used as a control (page 222, lines 28-29). The results of TaqManTM PCR are reported in ΔC_t units, as explained in the passage on page 222, lines 37-39. One unit corresponds to one PCR cycle or approximately a 2-fold amplification, relative to control, two units correspond to 4-fold, 3 units to 8-fold

amplification and so on. Using this PCR-based assay, Appellants showed that the gene encoding for PRO232 was amplified, that is, it showed approximately 1.04-2.40 Δ Ct units for five lung tumors and 1.00-2.41 Δ Ct units for seven colon tumors, which corresponds to $2^{1.04}$ - $2^{2.40}$ - fold amplification in lung or to $2^{1.00}$ - $2^{2.41}$ - fold amplification in colon tumors; that is **2.056-fold to 5.28-fold in five lung tumors** or **2.00-fold to 5.32-fold in seven colon tumors**, which would be considered significant and credible by one skilled in the art. Therefore, the PRO232 gene and the PRO232 polypeptide are important diagnostic markers to identify such malignant lung or colon cancers.

The Examiner has asserted that only five out of the nineteen lung cancer samples tested positive and only seven out of 17 colon tumor samples tested positive for the claimed polypeptide, thus, if a sample was taken from an individual with lung cancer for diagnosis, it is allegedly more likely than not that this assay would yield a false negative result. (Page 4 of the Office Action mailed December 27, 2007)

Appellants point out that, as any skilled artisan in the field of oncology would easily appreciate, not all tumor markers are generally associated with every tumor, or even, with most tumors. In fact, some tumor markers are useful for identifying rare malignancies. That is, the association of the tumor marker with a particular type of tumor lesion may be rare, or, the occurrence of that particular kind of tumor lesion itself may be rare. In either event, even these rare tumor markers, which may not give a positive hit with most common tumors, have great value in tumor diagnosis, and consequently, in tumor prognosis. For example, the article by Hanna and Mornin (submitted with the Response filed August 9, 2004), discloses that the known breast cancer marker HER-2/neu is “amplified and/or overexpressed in 10%-30% of invasive breast cancers and in 40%-60% of intraductal breast carcinoma” (page 1, col. 1). The skilled artisan would know that such tumor markers are very useful for better classification of tumors. Therefore, whether the PRO232 gene is amplified in most of the lung or colon tumors is not relevant to its identification as a tumor marker, or its patentable utility. Rather, whether the amplification data for PRO232 is significant is what lends support to its usefulness as a tumor marker.

It was well known in the art at the time of filing of the application that gene amplification, which occurs in most solid tumors like lung and colon cancers, is generally

associated with poor prognosis. Therefore, the PRO232 gene becomes an important diagnostic marker to identify such malignant lung and colon cancers, even if the malignancy associated with the PRO232 molecule is a rare occurrence and the present specification clearly discloses enough evidence such that, one skilled in the art would know that PRO232 is useful as a valuable diagnostic marker for identifying certain types of lung and colon cancers.

Appellants further note that the tumors listed in Table 8 are not similar tumors from different patients, but various types/classes of lung and/or colon tumors at different stages. Accordingly, a positive result from one tumor, where the nucleic acid was amplified, but not from other tumors, indicates that the nucleic acid can be used as a marker for diagnosing the presence of that kind of tumor in which it was amplified. Amplification of the nucleic acid would be indicative of that specific class of lung or colon tumor, whereas absence of amplification would be non-conclusive. The skilled artisan would certainly know that such tumor markers are useful for better classification of tumors. Therefore, whether the PRO232 gene is amplified in five lung tumors or in all lung tumors is not relevant to its identification as a tumor marker, or its patentable utility. The fact that not all lung tumors tested positive in this study does not make the gene amplification data less significant. Rather, the fact that the amplification data for PRO232 is considered significant is what lends support to its usefulness as a tumor marker. If the goal is to diagnose lung and colon cancers, then contrary to the Examiner's assertion, a positive result does indicate the presence of cancer, while a negative result is not conclusive, and requires follow up testing.

The Examiner has further alleged, based on Sen *et al.* and Hittelman *et al.*, that the observed gene amplification was not corrected for aneuploidy. The Examiner has also asserted it is not clear whether PRO232 is amplified in cancerous lung tissue more than in damaged lung tissue. (Pages 4-6 of the Office Action mailed December 27, 2007)

Appellants respectfully disagree and submit that their gene amplification data was not due to aneuploidy. Appellants have submitted the Ashkenazi Declaration to show that "detection of gene amplification can be used for cancer diagnosis even if the determination includes measurement of chromosomal aneuploidy." Regarding Sen *et al.* and Hittelman *et al.*, Appellants agree that while aneuploidy can be a feature of damaged tissue as well, besides cancerous or pre-cancerous tissue, and may not invariably lead to cancer, Sen *et al.* in fact

support the Appellants' position that PRO232 is still useful in diagnosing pre-cancerous lesions or cancer itself. For instance, the art in lung cancer at the time of filing of the instant application clearly described that "epithelial tumors develop through a multistep process driven by genetic instability" in damaged lung lesions which may eventually lead to lung cancer. Many articles published around June 23, 1999 (the effective filing date of this application) studied such damaged or premalignant lesions and suggested that identification of such pre-cancerous lesions were very important in preventive diagnosis and treatment of lung cancer. Based on the well-known art, Appellants submit that there is utility in identifying genetic biomarkers in epithelial tissues at cancer risk.

The Examiner has also addressed the pooled blood controls used in the gene amplification assay and asserted that the controls were not matched, non-tumor lung samples, but rather pooled DNA samples from blood of healthy subjects, citing Pennica *et al* and Konopka *et al.* as supporting references. (Page 10 of the Office Action mailed December 27, 2007)

Appellants respectfully submit that the Examiner's position is scientifically incorrect because the instant application relies on genomic DNA amplification for utility and not cDNA expression. Different types of cells from the same organism should have the same set of genomic DNA. Thus, it does not matter what kind of cells you use for the control as long as the control cells have the entire genome. Accordingly, a "tissue-matched" control is not necessary in the gene amplification assay.

Appellants further point out that Pennica *et al.* teaches the exact same "pooled normal blood controls" as that used in the instant gene amplification assay (for instance, see page 14718, column 1 and Figure 5 of Pennica *et al.*). Further, the references Bieche *et al.* and Pitti *et al.*, submitted as Exhibits F and G with the Goddard Declaration, also used "pooled normal blood controls" as control. For instance, in Pitti *et al.* the authors used the same quantitative TaqMan PCR assay and pooled normal blood controls described in the instant specification, to study gene amplification in lung and colon cancer of DcR3, a decoy receptor for Fas ligand. Pitti *et al.* analyzed DNA copy number "in genomic DNA from 35 primary lung and colon tumors, relative to pooled genomic DNA from peripheral blood leukocytes (PBL) of 10 healthy donors." (Page 701, col. 1). The authors also analyzed mRNA expression of DcR3 in primary tumor tissue sections and found tumor-specific expression, confirming the finding of frequent amplification in

tumors, and confirming that the pooled blood sample was a valid negative control for the gene amplification experiments. In Bieche et al., the authors used the quantitative TaqMan PCR assay to study gene amplification of *myc*, *ccnd1* and *erbB2* in breast tumors. As their negative control, Bieche et al. used normal leukocyte DNA derived from a small subset of the breast cancer patients (page 663). The authors note that "[t]he results of this study are consistent with those reported in the literature" (page 664, col. 2). Thus, contrary to the Examiner's allegations, Pennica et al, Pitti et al. and Bieche et al. in fact, confirm the validity of use of the "pooled blood control" as a negative controls, and indicate that this control was widely utilized in the art at the time of filing of the instant application.

C. A prima facie case of lack of utility has not been established

The Examiner has asserted, based on Pennica *et al.*, Konopka *et al.*, Godbout *et al.*, and Li *et al.* that there is a general lack of correlation between gene amplification and mRNA expression, and, thus, the data in Table 9 does not provide a basis for utility or enablement of the claimed polypeptides (Pages 4-7 of the Final Office Action mailed December 27, 2007).

As a preliminary matter, Appellants respectfully submit that it is not a legal requirement to establish that gene amplification "necessarily" results in increased expression at the mRNA and polypeptide levels or that polypeptide levels can be "accurately predicted." As discussed above, the evidentiary standard to be used throughout *ex parte* examination of a patent application is a preponderance of the totality of the evidence under consideration. Accordingly, Appellants submit that in order to overcome the presumption of truth that an assertion of utility by the applicant enjoys, the Examiner must establish that **it is more likely than not** that one of ordinary skill in the art would doubt the truth of the statement of utility. Therefore, it is not legally required that there be a "necessary" correlation between the data presented and the claimed subject matter. The law requires only that one skilled in the art should accept that such a correlation is **more likely than not to exist**. Appellants respectfully submit that when the proper evidentiary standard is applied, a correlation must be acknowledged. Only after the Examiner has made a proper prima facie showing of lack of utility, does the burden of rebuttal shift to the applicant. The Examiner failed to meet this evidentiary burden.

Pennica *et al.*

Appellants submit that Pennica *et al.* does not show a lack of correlation between gene (DNA) amplification and mRNA levels. According to the quoted statement from Pennica *et al.*, "WISP-1 gene amplification in human lung tumors showed a correlation between DNA amplification and over-expression, whereas overexpression of WISP-3 RNA was seen in the absence of DNA amplification. In contrast, WISP-2 DNA was amplified in lung tumors, but its mRNA expression was significantly reduced in the majority of tumors compared with expression in normal colonic mucosa from the same patient." From this, the Examiner correctly concludes that increased copy number does not *necessarily* result in increased polypeptide expression. The standard, however, is not absolute certainty. The fact that in the case of a specific class of closely related molecules there seemed to be no correlation with gene amplification and the level of mRNA/protein expression, does not establish that it is more likely than not, in general, that such correlation does not exist. The Examiner has not shown whether the lack or correlation observed for the family of WISP polypeptides is typical, or is merely a discrepancy, an exception to the rule of correlation. Indeed, the working hypothesis among those skilled in the art is that, if a gene is amplified in cancer, the encoded protein is likely to be expressed at an elevated level. In fact, as noted even in Pennica *et al.*, "[a]n analysis of *WISP*-1 gene amplification and expression in human lung tumors *showed a correlation between DNA amplification and over-expression . . .*" (Pennica *et al.*, page 14722, left column, first full paragraph, emphasis added).

Accordingly, Appellants respectfully submit that Pennica *et al.* teaches nothing conclusive regarding the absence of correlation between amplification of a gene and over-expression of the encoded WISP polypeptide. More importantly, the teaching of Pennica *et al.* is specific to *WISP* genes. Pennica *et al.* has no teaching whatsoever about the correlation of gene amplification and protein expression in general.

Konopka *et al.*

Regarding Konopka *et al.*, Applicants submit that the Examiner has completely misinterpreted the teachings in the cited reference. Contrary to the Examiner's assertions, Konopka *et al.* does not support the position that DNA amplification is not correlated with mRNA overexpression. Konopka *et al.* show only that, of the cell lines known to have increased abl protein expression, only one had amplification of the abl gene (page 4051, col. 1). This

result proves only that increased mRNA and protein expression levels can result from causes other than gene amplification. Konopka *et al.* do not demonstrate that when gene amplification does occur, it does not result in increased mRNA and protein expression levels, particularly given that the cell line with amplification of the abl gene did show increased abl mRNA and protein expression levels. Furthermore, Konopka *et al.* supports Appellants' position that mRNA levels correlate with protein levels. Konopka *et al.* state that "the 8-kb mRNA that encodes P210^{c-abl} was detected at a 10-fold higher level in SK-CML7bt-333 (Fig. 3A, +) than in SK-CML16Bt-1 (B, +), which **correlated** with the relative level of P210^{c-abl} detected in each cell line. Analysis of additional cell lines demonstrated that the level of 8-kb mRNA **directly correlated** with the level of P210^{c-abl} (Table 1)" (page 4050, col. 2, emphasis added).

Godbout *et al.*

Regarding Godbout, the Examiner has asserted that Godbout *et al.* teaches that "a number of studies suggest that co-amplified genes are only overexpressed if they provide a selective advantage to the cells in which they are amplified." The Examiner further asserts that Godbout teaches "[i]t is generally accepted that co-amplified genes are not over-expressed unless they provide a selective growth advantage to the cell." (Pages 6-7 of the Office Action mailed December 27, 2007).

Appellants have previously made of record three more recent references, published in 2002, by Orntoft *et al.*, Hyman *et al.*, and Pollack *et al.*, (made of record in Applicants' Response filed on August 9, 2004), which collectively teach that in general, gene amplification increases mRNA expression. Appellants submit that these more recent references must be acknowledged as more accurately reflecting the state of the art regarding the correlation between gene amplification and transcript expression than the references cited by Godbout *et al.*

Applicants further maintain that Godbout *et al.* report that "there is a good correlation with DDX1 gene copy number, DDX1 transcript levels, and DDX1 protein levels in all cell lines studied." Thus, in these cancer cell lines, DDX1 mRNA and protein levels are correlated.

Moreover, selective advantage to cell survival is not the only mechanism by which genes impact cancer. Mechanistic data is not a requirement for the utility requirement. Hence, this rejection is improper. Applicants respectfully submit that, as discussed above, Orntoft *et al.*,

Hyman *et al.*, and Pollack *et al.*, (of record), collectively teach that gene amplification increases mRNA expression for large numbers of genes, which have not been identified as being oncogenes or as conferring any selective growth advantage on tumor cells. Thus, the art of record clearly shows that there is no requirement that a polypeptide must be a known oncogene or a protein otherwise known to be associated with tumor growth, in order for amplification of the gene encoding the protein to correlate with increased protein expression. In fact, as demonstrated by Orntoft *et al.*, Hyman *et al.*, and Pollack *et al.*, examination of gene amplification is a useful way to identify novel proteins not previously known to be associated with cancer.

The Examiner has asserted that unlike Godbout *et al.*, the instant specification does not teach structure/ function analysis and the Examiner questions whether the level of genomic amplification of DDX1 gene is comparable to that disclosed by PRO232. (Page 12 of the Office Action mailed in December 27, 2007)

Appellants respectfully submit that it was never claimed that PRO232 is similar in any way to the DDX1 gene of Godbout *et al.*, they never claimed PRO232 was an RNA helicase or that it confers selective advantage to cell survival; on the other hand, the Godbout reference was submitted to show good correlation between protein levels based upon genomic DNA amplification, which the Examiner clearly agrees with. Moreover, selective advantage to cell survival is not the only mechanism by which genes impact cancer. Structure/function data, which the Examiner requests, is not a requirement for the utility requirement. Hence this rejection is improper.

Li *et al.*

The Examiner has cited Li *et al.* as teaching that “68.8% of the genes showing over-representation in the genome did not show elevated transcript levels.” (Page 8 of the Office Action mailed in December 27, 2007)

Appellants respectfully point out that Li *et al.* acknowledge that their results differed from those obtained by Hyman *et al.* and Pollack *et al.* (of record), who found a substantially higher level of correlation between gene amplification and increased gene expression. The authors note that “[t]his discordance may reflect methodologic differences between studies or biological differences between breast cancer and lung adenocarcinoma” (page 2629, col. 1). For

instance, as explained in the Supplemental Information accompanying the Li article, genes were considered to be amplified if they had a copy number ratio of at least 1.40. In the case of PRO830, as discussed in previously filed responses and in the Goddard Declaration (of record), an appropriate threshold for considering gene amplification to be significant is a copy number of at least 2.0 (which is a higher threshold). The PRO232 gene showed significant amplification of **2.0-fold to 5.2-fold** in twelve different lung and colon primary tumors, and thus fully meets this standard. It is not surprising that in the Li *et al.* reference, by using a lower threshold of 1.4 for considering gene amplification, a higher number of genes not showing corresponding increases in mRNA expression were found. Moreover, Appellants add that the results of Li *et al.* do not conclusively disprove that a gene with a substantially higher level of gene amplification, such as PRO232, would be expected to show a corresponding increase in transcript expression.

In response to Applicants' argument that the discordance may reflect methodologic differences, the Examiner asserts that "Li et al. did not limit their studies to genes that were amplified at less than 2-fold." In support of this assertion, the Examiner cites the first paragraph of the Supplemental Material. (Page 13 of the Office Action mailed in December 27, 2007)

Applicants respectfully point out that the Examiner has misinterpreted the methodology disclosed in the supplemental material. The evidence cited by the Examiner pertains to the inclusion criteria of the probes used for defining amplicons. In the second paragraph entitled "Relationship between genomic copy number and gene transcript level", the authors state that "[f]or each gene, the CGH data were represented by a vector that was labeled '1' for genomic overrepresentation (including amplification) ratio greater than 1.40 and '0' for no genomic overrepresentation." Nevertheless, the Examiner acknowledges that the alleged 2-fold amplification criteria would only apply to some of the samples. The Examiner has not established that a correlation does not exist in samples based solely on this threshold.

In summary, the Patent Office has failed to meet its initial burden of proof that Applicants' claims of utility are not substantial or credible. The arguments presented by the Examiner in combination with the cited articles do not provide sufficient reasons to doubt the statements by Applicants that PRO232 has utility. As discussed above, the law does not require that DNA amplification is "always" associated with overexpression of the gene product. Therefore, Applicants submit that the Examiner's reasoning is based on a misrepresentation of

the scientific data presented in the above cited reference and application of an improper, heightened legal standard. In fact, contrary to what the Examiner contends, the art indicates that, if a gene is amplified in cancer, it is more likely than not that the encoded protein will be expressed at an elevated level.

It is "more likely than not" for amplified genes to have increased mRNA

On the contrary, Appellants submit that Example 92 of the specification further discloses that, "(a)mplification is associated with overexpression of the gene product, indicating that the polypeptides are useful targets for therapeutic intervention in certain cancers such as lung, colon, breast and other cancers and diagnostic determination of the presence of those cancers" (Emphasis added). Besides, Appellants have submitted ample evidence to show that, in general, if a gene is amplified in cancer, it is "more likely than not" that the corresponding mRNA will also be expressed at an elevated level.

For instance, Appellants presented the articles by Orntoft *et al.*, Hyman *et al.*, and Pollack *et al.* (made of record in Appellants' Response filed August 9, 2004), who collectively teach that in general, for most genes, DNA amplification increases mRNA expression. The results presented by Orntoft *et al.*, Hyman *et al.*, and Pollack *et al.* are based upon wide ranging analyses of a large number of tumor associated genes. Orntoft *et al.* studied transcript levels of 5600 genes in malignant bladder cancers, many of which were linked to the gain or loss of chromosomal material, and found that in general (18 of 23 cases) chromosomal areas with more than 2-fold gain of DNA showed a corresponding increase in mRNA transcripts. Hyman *et al.* compared DNA copy numbers and mRNA expression of over 12,000 genes in breast cancer tumors and cell lines, and found that there was evidence of a prominent global influence of copy number changes on gene expression levels. In Pollack *et al.*, the authors profiled DNA copy number alteration across 6,691 mapped human genes in 44 predominantly advanced primary breast tumors and 10 breast cancer cell lines, and found that on average, a 2-fold change in DNA copy number was associated with a corresponding 1.5-fold change in mRNA levels. In summary, the evidence supports the Appellants' position that gene amplification is more likely than not predictive of increased mRNA and polypeptide levels.

Thus, the Examiner appears to disregard the ample evidence provided in the above referenced articles based on misinterpretations of their teachings. Appellants submit that in fact,

these articles lend significant support that for an amplified gene, it is more likely than not that the protein will also be overexpressed and would be viewed as reasonable and credible by one of ordinary skill in the art. The “more likely than not” standard is a much lower standard than a “necessary” correlation or “accurate” prediction, and is clearly met in the claimed invention. Moreover, the Examiner has not cited any evidence or advanced any arguments as to why Appellants’ statement of overexpression of protein would not be credible. Accordingly, this point is believed to be moot.

Even if a *prima facie* case of lack of utility has been established, it should be withdrawn on consideration of the totality of evidence

Even if one assumes *arguendo* that it is more likely than not that there is no correlation between gene amplification and increased mRNA/protein expression, which Appellants submit is **not** true, a polypeptide encoded by a gene that is amplified in cancer would **still** have a specific, substantial, and credible utility. In support, Appellants respectfully draw the Board’s attention to page 2 of the Declaration of Dr. Avi Ashkenazi (submitted with the Response filed December 10, 2003) which explains that,

even when amplification of a cancer marker gene does not result in significant over-expression of the corresponding gene product, this very absence of gene product over-expression still provides significant information for cancer diagnosis and treatment. Thus, if over-expression of the gene product does not parallel gene amplification in certain tumor types but does so in others, then parallel monitoring of gene amplification and gene product over-expression enables more accurate tumor classification and hence better determination of suitable therapy. In addition, absence of over-expression is crucial information for the practicing clinician. If a gene is amplified but the corresponding gene product is not over-expressed, the clinician accordingly will decide not to treat a patient with agents that target that gene product.

Appellants thus submit that simultaneous testing of gene amplification and gene product over-expression enables more accurate tumor classification, even if the gene-product, the protein, is not over-expressed. This leads to better determination of a suitable therapy. Further, as explained in Dr. Ashkenazi’s Declaration, absence of over-expression of the protein itself is crucial information for the practicing clinician. If a gene is amplified in a tumor, but the

corresponding gene product is not over-expressed, the clinician will decide not to treat a patient with agents that target that gene product. This not only saves money, but also has the benefit that the patient can avoid exposure to the side effects associated with such agents.

This utility is further supported by the teachings of the article by Hanna and Mornin. (Pathology Associates Medical Laboratories, August (1999), submitted with the Response filed August 9, 2004). The article teaches that the HER-2/neu gene has been shown to be amplified and/or over-expressed in 10%-30% of invasive breast cancers and in 40%-60% of intraductal breast carcinomas. Further, the article teaches that diagnosis of breast cancer includes testing both the amplification of the HER-2/neu gene (by FISH) as well as the over-expression of the HER-2/neu gene product (by IHC). Even when the protein is not over-expressed, the assay relying on both tests leads to a more accurate classification of the cancer and a more effective treatment of it.

Appellants have clearly shown that the gene encoding the PRO232 polypeptide is amplified in at least five lung tumors and seven colon tumors. Therefore, the PRO232 gene, similar to the HER-2/neu gene disclosed in Hanna *et al.*, is a tumor associated gene. Furthermore, as discussed above, in the majority of amplified genes, the teachings in the art overwhelmingly show that gene amplification influences gene expression at the mRNA and protein levels. Therefore, one of skill in the art would reasonably expect in this instance, based on the amplification data for the PRO232 gene, that the PRO232 polypeptide is concomitantly overexpressed.

Thus, based on the asserted utility for PRO232 in the diagnosis of selected lung or colon tumors, the reduction to practice of the instantly claimed protein sequence of SEQ ID NO: 18 in the present application (also see pages 3- 4 and Example 4, pages 149-150), the step-by-step preparation of chimeric PRO polypeptides, including those wherein the heterologous polypeptide is an epitope tag or an Fc region of an immunoglobulin (page 74, line 23 to page 75, line 5), the description of the expression of PRO polypeptides in various host cells, including *E. coli*, mammalian cells, yeast and Baculovirus-infected insect cells at least in Examples 53-56, pages 192-199, the disclosure of the step-by-step protocol for the preparation, isolation and detection of monoclonal, polyclonal and other types of antibodies against the PRO232 protein in the specification (monoclonal and polyclonal antibodies at page 139, line 32, to page 141,

line 13; humanized antibodies at page 141, line 15, to page 142, line 16; antibody fragments at page 143, line 8 onwards; labeled antibodies at pages 144-145, line 16 onwards and page 146, line 33 to page 147, line 6) and the disclosure of the gene amplification assay in Example 92, the skilled artisan would know exactly how to make and use the claimed polypeptides for the diagnosis of lung or colon cancers. Appellants submit that based on the detailed information presented in the specification and the advanced state of the art in oncology, the skilled artisan would have found such testing routine and not 'undue.'

Therefore, since the instantly claimed invention is supported by either a credible, specific and substantial asserted utility or a well-established utility, one skilled in the art would know "how to make and use" the claimed invention without undue experimentation, Appellants respectfully request reconsideration and reversal of the determination of priority for Claims 44-46 and 49-51.

ISSUE 2. Claims 44-46 and 49-51 should be entitled to the priority date of U.S. Provisional Patent Application Serial No. 60/059121, filed September 17, 1997

The instant application has not been granted the earlier priority date of U.S. Provisional Patent Application Serial No. 60/059121, filed September 17, 1997 on the grounds that the prior 60/059121 application fails to provide a utility and lacks an enabling disclosure for the claimed invention under 35 U.S.C. §§101/112, first paragraph."

Appellants disagree and submit that, for the same reasons discussed above under Issue 1, U.S. Provisional Patent Application Serial No. 60/059121 also satisfies the utility requirements. Therefore, Appellants should be entitled to the priority date of **September 17, 1997**.

ISSUE 3. Claims 44-46 and 49-51 are not anticipated by Rosenthal *et al.*, DE19818619-A1 (dated 10/1999)

Claims 44-46 and 49-51 remain rejected under 35 U.S.C. §102(b) as being anticipated by Rosenthal *et al.*, DE19818619-A1 (dated 10/1999).

For the reasons discussed above under Issue 2, Appellants maintain that they are entitled to an effective filing date of September 17, 1997 based on a properly claimed priority to International application PCT/US98/18824. Therefore, Rosenthal *et al.* is not prior art and does not anticipate the instant claims. Accordingly, this rejection under 35 U.S.C. §102(b) should be withdrawn.

CONCLUSION

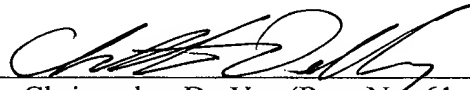
For the reasons given above, Appellants submit that present specification and the specification of U.S. Provisional Patent Application Serial No. 60/059121 dated September 17, 1997 clearly describes and provides at least one patentable utility for the instantly claimed invention. Moreover, it is respectfully submitted that the present specification clearly teaches "how to use" the presently claimed polypeptide based upon this disclosed patentable utility. Accordingly, Rosenthal *et al.*, DE19818619-A1 is not prior art. As such, Appellants respectfully request reconsideration and reversal of the outstanding rejection of Claims 44-46 and 49-51.

The Commissioner is authorized to charge any fees which may be required, including extension fees, or credit any overpayment to Deposit Account No. **50-4634** (referencing Attorney's Docket No. **123851-181890 (GNE-1618P2C18)**).

Respectfully submitted,

Date: September 29, 2008

By:


Christopher De Vry (Reg. No. 61,425)

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VIII. CLAIMS APPENDIX

Claims on Appeal

44. An isolated polypeptide comprising:
- (a) the amino acid sequence of the polypeptide of SEQ ID NO:18;
 - (b) the amino acid sequence of the polypeptide of SEQ ID NO:18, lacking its associated signal peptide; or
 - (c) the amino acid sequence of the polypeptide encoded by the full-length coding sequence of the cDNA deposited under ATCC accession number 209250;
- wherein, the nucleic acid encoding said polypeptide is amplified in lung or colon tumors.
45. The isolated polypeptide of Claim 44 comprising the amino acid sequence of the polypeptide of SEQ ID NO:18.
46. The isolated polypeptide of Claim 44 comprising the amino acid sequence of the polypeptide of SEQ ID NO:18, lacking its associated signal peptide.
49. The isolated polypeptide of Claim 44 comprising the amino acid sequence of the polypeptide encoded by the full-length coding sequence of the cDNA deposited under ATCC accession number 209250.
50. A chimeric polypeptide comprising a polypeptide according to Claim 44 fused to a heterologous polypeptide.
51. The chimeric polypeptide of Claim 50, wherein said heterologous polypeptide is an epitope tag or an Fc region of an immunoglobulin.

IX. EVIDENCE APPENDIX

1. Declaration of Avi Ashkenazi, Ph.D. under 35 C.F.R. §1.132, with attached Exhibit A (Curriculum Vitae).
2. Declaration of Audrey D. Goddard, Ph.D. under 37 C.F.R. §1.132, with attached Exhibits A-G:
 - A. Curriculum Vitae of Audrey D. Goddard, Ph.D.
 - B. Higuchi, R. *et al.*, "Simultaneous amplification and detection of specific DNA sequences," *Biotechnology* 10:413-417 (1992).
 - C. Livak, K.J., *et al.*, "Oligonucleotides with fluorescent dyes at opposite ends provide a quenched probe system useful for detecting PCR product and nucleic acid hybridization," *PCR Methods Appl.* 4:357-362 (1995).
 - D. Heid, C.A. *et al.*, "Real time quantitative PCR," *Genome Res.* 6:986-994 (1996).
 - E. Pennica, D. *et al.*, "WISP genes are members of the connective tissue growth factor family that are up-regulated in Wnt-1-transformed cells and aberrantly expressed in human colon tumors," *Proc. Natl. Acad. Sci. USA* 95:14717-14722 (1998).
 - F. Pitti, R.M. *et al.*, "Genomic amplification of a decoy receptor for Fas ligand in lung and colon cancer," *Nature* 396:699-703 (1998).
 - G. Bieche, I. *et al.*, "Novel approach to quantitative polymerase chain reaction using real-time detection: Application to the detection of gene amplification in breast cancer," *Int. J. Cancer* 78:661-666 (1998).
3. Declaration of Paul Polakis, Ph.D. under 37 C.F.R. §1.132.
4. Hyman, E., *et al.*, "Impact of DNA Amplification on Gene Expression Patterns in Breast Cancer," *Cancer Research* 62:6240-6245 (2002).
5. Pollack, J.R., *et al.*, "Microarray Analysis Reveals a Major Direct Role of DNA Copy Number Alteration in the Transcriptional Program of Human Breast Tumors," *Proc. Natl. Acad. Sci. USA* 99:12963-12968 (2002).
6. Orntoft, T.F., *et al.* *Molecular & Cellular Proteomics* – 1:37-45 (2002).
7. Hanna *et al.*, "HER-2/neu Breast Cancer Predictive Testing," Pathology Associates Medical Laboratories (1999).
8. Declaration of Paul Polakis, Ph.D. under 35 C.F.R. §1.132 (Polakis II).
9. Godbout, R., *et al.*, *J Biol Chem.* - 273(33):21161-8 (1998).
10. Rosenthal *et al.*, DE19818619-A1 (dated October 28, 1999).

11. Sen, S., "Aneuploidy and cancer," *Curr. Opin. Oncol.* **12**:82-88 (2000).
12. Li *et al.*, 2006, *Oncogene*, **25**: 2628-2635.
13. Konopka *et al.*, "Variable Expression of the Translocated c-abl oncogene in Philadelphia-chromosome-positive B-lymphoid cell lines from chronic myelogenous leukemia patients" *Proc. Natl. Acad. Sci. USA* **83**: 4049-52, (1986).
14. Hittelman, 2001, *Ann. N. Y. Acad. Sci.* **952**:1-12.

Item 1 was submitted with Appellants' Response filed December 10, 2003, and was considered by the Examiner as indicated in the Office action mailed April 8, 2004.

Item 2 was submitted with Appellants' Response filed July 25, 2005, and were considered by the Examiner as indicated in the second Final Office action mailed October 21, 2005.

Items 3-7 were submitted with Appellants' Response filed August 9, 2004, and were considered by the Examiner as indicated in the Office action mailed December 6, 2004.

Items 8 and were submitted with Appellants' Response filed July 27, 2006, and was considered by the Examiner as indicated in the Final Office Action mailed October 17, 2006.

Item 9 was submitted with Appellants' Appeal Brief filed September 24, 2007 and made of records by the Examiner in the Final Office Action mailed December 27, 2007.

Item 1 (E), *i.e.*: Pennica *et al.*, and Items 10-11 were made of record by the Examiner in the Office Action mailed September 29, 2003.

Item 12 was made of record by the Examiner in the Final Office Action mailed October 17, 2006.

Items 13 and 14 were made of record by the Examiner in the Final Office Action mailed December 27, 2007.

X. RELATED PROCEEDINGS APPENDIX

None- no decision rendered by a Court or the Board in any related proceedings identified above.

LIBC/3392711.1

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Ashkenazi et al.	Group Art Unit: 1647
Serial No.: 09/903,925	Examiner: Fozia Hamid
Filed: July 11, 2001	CERTIFICATE OF MAILING I hereby certify that this correspondence is being deposited with the United States Postal Service with sufficient postage as first class mail in an envelope addressed to: Assistant Commissioner of Patents, Washington, D.C. 20231 on Date: _____
For: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC ACIDS	

DECLARATION OF AUDREY D. GODDARD, Ph.D UNDER 37 C.F.R. § 1.132

Assistant Commissioner of Patents
Washington, D.C. 20231

Sir:

I, Audrey D. Goddard, Ph.D. do hereby declare and say as follows:

1. I am a Senior Clinical Scientist at the Experimental Medicine/BioOncology, Medical Affairs Department of Genentech, Inc., South San Francisco, California 94080.
2. Between 1993 and 2001, I headed the DNA Sequencing Laboratory at the Molecular Biology Department of Genentech, Inc. During this time, my responsibilities included the identification and characterization of genes contributing to the oncogenic process, and determination of the chromosomal localization of novel genes.
3. My scientific Curriculum Vitae, including my list of publications, is attached to and forms part of this Declaration (Exhibit A).

Serial No.: *

Filed: *

4. I am familiar with a variety of techniques known in the art for detecting and quantifying the amplification of oncogenes in cancer, including the quantitative TaqMan PCR (i.e., "gene amplification") assay described in the above captioned patent application.

5. The TaqMan PCR assay is described, for example, in the following scientific publications: Higuchi *et al.*, Biotechnology 10:413-417 (1992) (Exhibit B); Livak *et al.*, PCR Methods Appl., 4:357-362 (1995) (Exhibit C) and Heid *et al.*, Genome Res. 6:986-994 (1996) (Exhibit D). Briefly, the assay is based on the principle that successful PCR yields a fluorescent signal due to Taq DNA polymerase-mediated exonuclease digestion of a fluorescently labeled oligonucleotide that is homologous to a sequence between two PCR primers. The extent of digestion depends directly on the amount of PCR, and can be quantified accurately by measuring the increment in fluorescence that results from decreased energy transfer. This is an extremely sensitive technique, which allows detection in the exponential phase of the PCR reaction and, as a result, leads to accurate determination of gene copy number.

6. The quantitative fluorescent TaqMan PCR assay has been extensively and successfully used to characterize genes involved in cancer development and progression. Amplification of protooncogenes has been studied in a variety of human tumors, and is widely considered as having etiological, diagnostic and prognostic significance. This use of the quantitative TaqMan PCR assay is exemplified by the following scientific publications: Pennica *et al.*, Proc. Natl. Acad. Sci. USA 95(25):14717-14722 (1998) (Exhibit E); Pitti *et al.*, Nature 396(6712):699-703 (1998) (Exhibit F) and Bieche *et al.*, Int. J. Cancer 78:661-666 (1998) (Exhibit G), the first two of which I am co-author. In particular, Pennica *et al.* have used the quantitative TaqMan PCR assay to study relative gene amplification of WISP and c-myc in various cell lines, colorectal tumors and normal mucosa. Pitti *et al.* studied the genomic amplification of a decoy receptor for Fas ligand in lung and colon cancer, using the quantitative TaqMan PCR assay. Bieche *et al.* used the assay to study gene amplification in breast cancer.

Serial No.: *

Filed: *

7. It is my personal experience that the quantitative TaqMan PCR technique is technically sensitive enough to detect at least a 2-fold increase in gene copy number relative to control. It is further my considered scientific opinion that an at least 2-fold increase in gene copy number in a tumor tissue sample relative to a normal (i.e., non-tumor) sample is significant and useful in that the detected increase in gene copy number in the tumor sample relative to the normal sample serves as a basis for using relative gene copy number as quantitated by the TaqMan PCR technique as a diagnostic marker for the presence or absence of tumor in a tissue sample of unknown pathology. Accordingly, a gene identified as being amplified at least 2-fold by the quantitative TaqMan PCR assay in a tumor sample relative to a normal sample is useful as a marker for the diagnosis of cancer, for monitoring cancer development and/or for measuring the efficacy of cancer therapy.

8. I declare further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true. I declare that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Jan. 16, 2003

Date

Audrey D. Goddard

Audrey D. Goddard, Ph.D.

AUDREY D. GODDARD, Ph.D.

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1 DNA Way
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650.225.6429
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110 Congo St.
San Francisco, CA, 94131
415.841.9154
415.819.2247 (mobile)
agoddard@pacbell.net

PROFESSIONAL EXPERIENCE

Genentech, Inc.
South San Francisco, CA

1993-present

2001 - present Senior Clinical Scientist
Experimental Medicine / BioOncology, Medical Affairs

Responsibilities:

- *Companion diagnostic oncology products*
- *Acquisition of clinical samples from Genentech's clinical trials for translational research*
- *Translational research using clinical specimen and data for drug development and diagnostics*
- *Member of Development Science Review Committee, Diagnostic Oversight Team, 21 CFR Part 11 Subteam*

Interests:

- *Ethical and legal implications of experiments with clinical specimens and data*
- *Application of pharmacogenomics in clinical trials*

1998 - 2001 Senior Scientist

Head of the DNA Sequencing Laboratory, Molecular Biology Department, Research

Responsibilities:

- *Management of a laboratory of up to nineteen –including postdoctoral fellow, associate scientist, senior research associate and research assistants/associate levels*
- *Management of a \$750K budget*
- *DNA sequencing core facility supporting a 350+ person research facility.*
- *DNA sequencing for high throughput gene discovery, - ESTs, cDNAs, and constructs*
- *Genomic sequence analysis and gene identification*
- *DNA sequence and primary protein analysis*

Research:

- *Chromosomal localization of novel genes*
- *Identification and characterization of genes contributing to the oncogenic process*
- *Identification and characterization of genes contributing to inflammatory diseases*
- *Design and development of schemes for high throughput genomic DNA sequence analysis*
- *Candidate gene prediction and evaluation*

1993 - 1998 Scientist

Head of the DNA Sequencing Laboratory, Molecular Biology Department, Research

Responsibilities

- *DNA sequencing core facility supporting a 350+ person research facility*
- *Assumed responsibility for a pre-existing team of five technicians and expanded the group into fifteen, introducing a level of middle management and additional areas of research*
- *Participated in the development of the basic plan for high throughput secreted protein discovery program – sequencing strategies, data analysis and tracking, database design*
- *High throughput EST and cDNA sequencing for new gene identification.*
- *Design and implementation of analysis tools required for high throughput gene identification.*
- *Chromosomal localization of genes encoding novel secreted proteins.*

Research:

- *Genomic sequence scanning for new gene discovery.*
- *Development of signal peptide selection methods.*
- *Evaluation of candidate disease genes.*
- *Growth hormone receptor gene SNPs in children with Idiopathic short stature*

**Imperial Cancer Research Fund
London, UK with Dr. Ellen Solomon**

1989-1992

6/89 -12/92 Postdoctoral Fellow

- Cloning and characterization of the genes fused at the acute promyelocytic leukemia translocation breakpoints on chromosomes 17 and 15.
- Prepared a successfully funded European Union multi-center grant application

**McMaster University
Hamilton, Ontario, Canada with Dr. G. D. Sweeney**

1983

5/83 – 8/83: NSERC Summer Student

- *In vitro* metabolism of β -naphthoflavone in C57BL/6J and DBA mice

EDUCATION

Ph.D.

"Phenotypic and genotypic effects of mutations in the human retinoblastoma gene."

Supervisor: Dr. R. A. Phillips

University of Toronto
Toronto, Ontario, Canada.
Department of Medical
Biophysics.

1989

Honours B.Sc

"The *in vitro* metabolism of the cytochrome P-448 inducer β -naphthoflavone in C57BL/6J mice."

Supervisor: Dr. G. D. Sweeney

McMaster University,
Hamilton, Ontario, Canada.
Department of Biochemistry

1983

ACADEMIC AWARDS

Imperial Cancer Research Fund Postdoctoral Fellowship	1989-1992
Medical Research Council Studentship	1983-1988
NSERC Undergraduate Summer Research Award	1983
Society of Chemical Industry Merit Award (Hons. Biochem.)	1983
Dr. Harry Lyman Hooker Scholarship	1981-1983
J.L.W. Gill Scholarship	1981-1982
Business and Professional Women's Club Scholarship	1980-1981
Wyerhauser Foundation Scholarship	1979-1980

INVITED PRESENTATIONS

Genentech's gene discovery pipeline: High throughput identification, cloning and characterization of novel genes. Functional Genomics: From Genome to Function, Litchfield Park, AZ, USA. October 2000

High throughput identification, cloning and characterization of novel genes. G2K:Back to Science, Advances in Genome Biology and Technology I. Marco Island, FL, USA. February 2000

Quality control in DNA Sequencing: The use of Phred and Phrap. Bay Area Sequencing Users Meeting, Berkeley, CA, USA. April 1999

High throughput secreted protein identification and cloning. Tenth International Genome Sequencing and Analysis Conference, Miami, FL, USA. September 1998

The evolution of DNA sequencing: The Genentech perspective. Bay Area Sequencing Users Meeting, Berkeley, CA, USA. May 1998

Partial Growth Hormone Insensitivity: The role of GH-receptor mutations in Idiopathic Short Stature. Tenth Annual National Cooperative Growth Study Investigators Meeting, San Francisco, CA, USA. October, 1996

Growth hormone (GH) receptor defects are present in selected children with non-GH-deficient short stature: A molecular basis for partial GH-insensitivity. 76th Annual Meeting of The Endocrine Society, Anaheim, CA, USA. June 1994

A previously uncharacterized gene, myl, is fused to the retinoic acid receptor alpha gene in acute promyelocytic leukemia. XV International Association for Comparative Research on Leukemia and Related Disease, Padua, Italy. October 1991

PATENTS

Goddard A, Godowski PJ, Gurney AL. NL2 Tie ligand homologue polypeptide. Patent Number: 6,455,496. Date of Patent: Sept. 24, 2002.

Goddard A, Godowski PJ and Gurney AL. NL3 Tie ligand homologue nucleic acids. Patent Number: 6,426,218. Date of Patent: July 30, 2002.

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RESEARCH

SIMULTANEOUS AMPLIFICATION AND DETECTION OF SPECIFIC DNA SEQUENCES

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We have enhanced the polymerase chain reaction (PCR) such that specific DNA sequences can be detected without opening the reaction tube. This enhancement requires the addition of ethidium bromide (EtBr) to a PCR. Since the fluorescence of EtBr increases in the presence of double-stranded (ds) DNA an increase in fluorescence in such a PCR indicates a positive amplification, which can be easily monitored externally. In fact, amplification can be continuously monitored in order to follow its progress. The ability to simultaneously amplify specific DNA sequences and detect the product of the amplification both simplifies and improves PCR and may facilitate its automation and more widespread use in the clinic or in other situations requiring high sample throughput.

Although the potential benefits of PCR¹ to clinical diagnostics are well known^{2,3}, it is still not widely used in this setting, even though it is four years since thermostable DNA polymerases⁴ made PCR practical. Some of the reasons for its slow acceptance are high cost, lack of automation of pre- and post-PCR processing steps, and false positive results from carryover-contamination. The first two points are related in that labor is the largest contributor to cost at the present stage of PCR development. Most current assays require some form of "downstream" processing once thermocycling is done in order to determine whether the target DNA sequence was present and has amplified. These include DNA hybridization^{5,6}, gel electrophoresis with or without use of restriction digestion^{7,8}, HPLC⁹, or capillary electrophoresis¹⁰. These methods are labor-intensive, have low throughput, and are difficult to automate. The third point is also closely related to downstream processing. The handling of the PCR product in these downstream processes increases the chances that amplified DNA will spread through the typing lab, resulting in a risk of

"carryover" false positives in subsequent testing¹¹.

These downstream processing steps would be eliminated if specific amplification and detection of amplified DNA took place simultaneously within an unopened reaction vessel. Assays in which such different processes take place without the need to separate reaction components have been termed "homogeneous". No truly homogeneous PCR assay has been demonstrated to date, although progress towards this end has been reported. Chehab, et al.¹², developed a PCR product detection scheme using fluorescent primers that resulted in a fluorescent PCR product. Allele-specific primers, each with different fluorescent tags, were used to indicate the genotype of the DNA. However, the unincorporated primers must still be removed in a downstream process in order to visualize the result. Recently, Holland, et al.¹³, developed an assay in which the endogenous 5' exonuclease assay of *Taq* DNA polymerase was exploited to cleave a labeled oligonucleotide probe. The probe would only cleave if PCR amplification had produced its complementary sequence. In order to detect the cleavage products, however, a subsequent process is again needed.

We have developed a truly homogeneous assay for PCR and PCR product detection based upon the greatly increased fluorescence that ethidium bromide and other DNA binding dyes exhibit when they are bound to dsDNA¹⁴⁻¹⁶. As outlined in Figure 1, a prototypic PCR

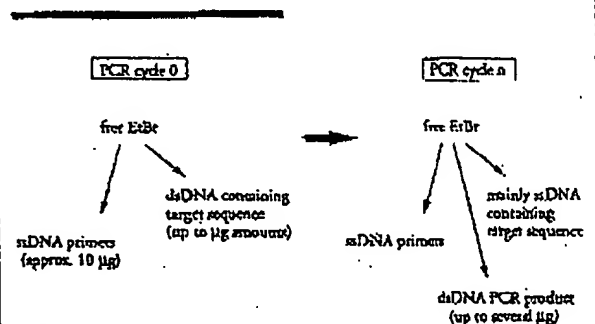


FIGURE 1 Principle of simultaneous amplification and detection of PCR product. The components of a PCR containing EtBr that are fluorescent are listed—EtBr itself, EtBr bound to either ssDNA or dsDNA. There is a large fluorescence enhancement when EtBr is bound to DNA and binding is greatly enhanced when DNA is double-stranded. After sufficient (n) cycles of PCR, the net increase in dsDNA results in additional EtBr binding, and a net increase in total fluorescence.

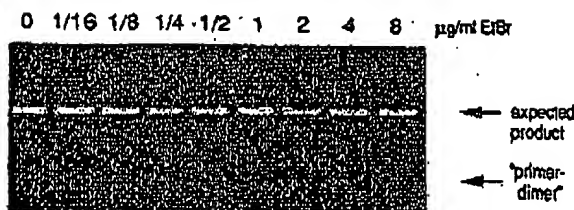


FIGURE 2 Gel electrophoresis of PCR amplification products of the human nuclear gene, HLA DQα, made in the presence of increasing amounts of EtBr (up to 8 μg/ml). The presence of EtBr has no obvious effect on the yield or specificity of amplification.

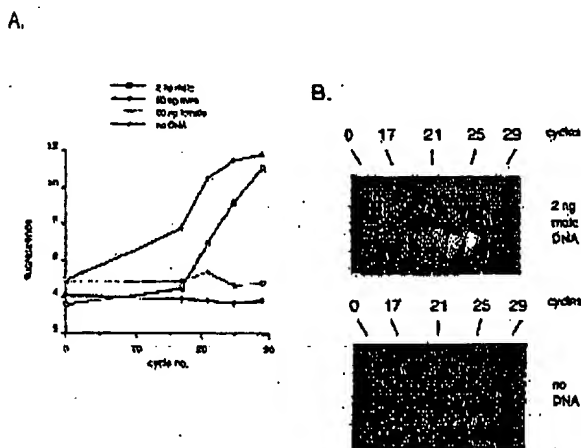


FIGURE 3 (A) Fluorescence measurements from PCR reactions that contain 0.5 μg/ml EtBr and that are specific for Y-chromosome repeat sequences. Five replicate PCRs were begun containing each of the DNAs specified. At each indicated cycle, one of the five replicate PCRs for each DNA was removed from thermocycling and its fluorescence measured. Units of fluorescence are arbitrary. (B) UV photograph of PCR tubes (0.5 ml Eppendorf-style, polypropylene micro-centrifuge tubes) containing reactions, those starting from 2 ng male DNA and control reactions without any DNA, from (A).

begins with primers that are single-stranded DNA (ssDNA), dNTPs, and DNA polymerase. An amount of dsDNA containing the target sequence (target DNA) is also typically present. This amount can vary, depending on the application, from single-cell amounts of DNA¹⁷ to micrograms per PCR¹⁸. If EtBr is present, the reagents that will fluoresce, in order of increasing fluorescence, are free EtBr itself, and EtBr bound to the single-stranded DNA primers and to the double-stranded target DNA (by its intercalation between the stacked bases of the DNA double-helix). After the first denaturation cycle, target DNA will be largely single-stranded. After a PCR is completed, the most significant change is the increase in the amount of dsDNA (the PCR product itself) of up to several micrograms. Formerly free EtBr is bound to the additional dsDNA, resulting in an increase in fluorescence. There is also some decrease in the amount of ssDNA primer, but because the binding of EtBr to ssDNA is much less than to dsDNA, the effect of this change on the total fluorescence of the sample is small. The fluorescence increase can be measured by directing excitation illumination through the walls of the amplification vessel

before and after, or even continuously during, thermocycling.

RESULTS

PCR in the presence of EtBr. In order to assess the effect of EtBr in PCR, amplifications of the human HLA DQα gene¹⁹ were performed with the dye present at concentrations from 0.06 to 8.0 μg/ml (a typical concentration of EtBr used in staining of nucleic acids following gel electrophoresis is 0.5 μg/ml). As shown in Figure 2, gel electrophoresis revealed little or no difference in the yield or quality of the amplification product whether EtBr was absent or present at any of these concentrations, indicating that EtBr does not inhibit PCR.

Detection of human Y-chromosome specific sequences. Sequence-specific fluorescence enhancement of EtBr as a result of PCR was demonstrated in a series of amplifications containing 0.5 μg/ml EtBr and primers specific to repeat DNA sequences found on the human Y-chromosome²⁰. These PCRs initially contained either 60 ng male, 60 ng female, 2 ng roak human or no DNA. Five replicate PCRs were begun for each DNA. After 0, 17, 21, 24 and 29 cycles of thermocycling, a PCR for each DNA was removed from the thermocycler, and its fluorescence measured in a spectrofluorometer and plotted vs. amplification cycle number (Fig. 3A). The shape of this curve reflects the fact that by the time an increase in fluorescence can be detected, the increase in DNA is becoming linear and not exponential with cycle number. As shown, the fluorescence increased about three-fold over the background fluorescence for the PCRs containing human male DNA, but did not significantly increase for negative control PCRs, which contained either no DNA or human female DNA. The more male DNA present to begin with—60 ng versus 2 ng—the fewer cycles were needed to give a detectable increase in fluorescence. Gel electrophoresis on the products of these amplifications showed that DNA fragments of the expected size were made in the male DNA containing reactions and that little DNA synthesis took place in the control samples.

In addition, the increase in fluorescence was visualized by simply laying the completed, unopened PCRs on a UV transilluminator and photographing them through a red filter. This is shown in figure 3B for the reactions that began with 2 ng male DNA and those with no DNA.

Detection of specific alleles of the human β-globin gene. In order to demonstrate that this approach has adequate specificity to allow genetic screening, a detection of the sickle-cell anemia mutation was performed. Figure 4 shows the fluorescence from completed amplifications containing EtBr (0.5 μg/ml) as detected by photography of the reaction tubes on a UV transilluminator. These reactions were performed using primers specific for either the wild-type or sickle-cell mutation of the human β-globin gene²¹. The specificity for each allele is imparted by placing the sickle-mutation site at the terminal 3' nucleotide of one primer. By using an appropriate primer annealing temperature, primer extension—and thus amplification—can take place only if the 3' nucleotide of the primer is complementary to the β-globin allele present²².

Each pair of amplifications shown in Figure 4 consists of a reaction with either the wild-type allele specific (left tube) or sickle-allele specific (right tube) primers. Three different DNAs were typed: DNA from a homozygous, wild-type β-globin individual (AA); from a heterozygous sickle β-globin individual (AS); and from a homozygous sickle β-globin individual (SS). Each DNA (50 ng genomic DNA to start each PCR) was analyzed in triplicate (3 pairs

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of reactions each). The DNA type was reflected in the relative fluorescence intensities in each pair of completed amplifications. There was a significant increase in fluorescence only where a β -globin allele DNA matched the primer set. When measured on a spectrofluorometer (data not shown), this fluorescence was about three times that present in a PCR where both β -globin alleles were mismatched to the primer set. Gel electrophoresis (not shown) established that this increase in fluorescence was due to the synthesis of nearly a microgram of a DNA fragment of the expected size for β -globin. There was little synthesis of dsDNA in reactions in which the allele-specific primer was mismatched to both alleles.

Continuous monitoring of a PCR. Using a fiber optic device, it is possible to direct excitation illumination from a spectrofluorometer to a PCR undergoing thermocycling and to return its fluorescence to the spectrofluorometer. The fluorescence readout of such an arrangement, directed at an EtBr-containing amplification of Y-chromosome specific sequences from 25 ng of human male DNA, is shown in Figure 5. The readout from a control PCR with no target DNA is also shown. Thirty cycles of PCR were monitored for each.

The fluorescence trace as a function of time clearly shows the effect of the thermocycling. Fluorescence intensity rises and falls inversely with temperature. The fluorescence intensity is minimum at the denaturation temperature (94°C) and maximum at the annealing/extension temperature (50°C). In the negative-control PCR, these fluorescence maxima and minima do not change significantly over the thirty thermocycles, indicating that there is little dsDNA synthesis without the appropriate target DNA, and there is little if any bleaching of EtBr during the continuous illumination of the sample.

In the PCR containing male DNA, the fluorescence maxima at the annealing/extension temperature begin to increase at about 4000 seconds of thermocycling, and continue to increase with time, indicating that dsDNA is being produced at a detectable level. Note that the fluorescence minima at the denaturation temperature do not significantly increase, presumably because at this temperature there is no dsDNA for EtBr to bind. Thus the course of the amplification is followed by tracking the fluorescence increase at the annealing temperature. Analysis of the products of these two amplifications by gel electrophoresis showed a DNA fragment of the expected size for the male DNA containing sample and no detectable DNA synthesis for the control sample.

DISCUSSION

Downstream processes such as hybridization to a sequence-specific probe can enhance the specificity of DNA detection by PCR. The elimination of these processes means that the specificity of this homogeneous assay depends solely on that of PCR. In the case of sickle-cell disease, we have shown that PCR alone has sufficient DNA sequence specificity to permit genetic screening. Using appropriate amplification conditions, there is little non-specific production of dsDNA in the absence of the appropriate target allele.

The specificity required to detect pathogens can be more or less than that required to do genetic screening, depending on the number of pathogens in the sample and the amount of other DNA that must be taken with the sample. A difficult target is HIV, which requires detection of a viral genome that can be at the level of a few copies per thousands of host cells⁶. Compared with genetic screening, which is performed on cells containing at least one copy of the target sequence, HIV detection requires both more specificity and the input of more total

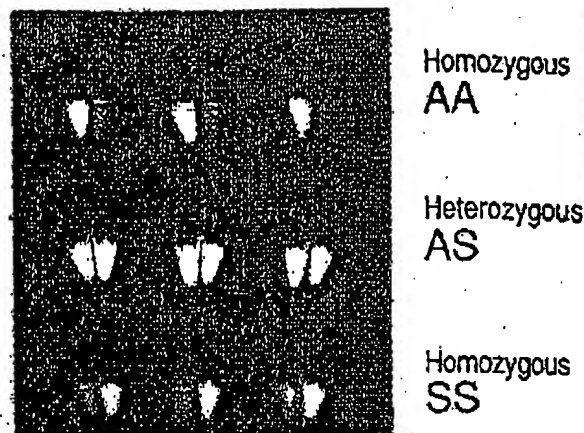


FIGURE 4 UV photograph of PCR tubes containing amplifications using EtBr that are specific to wild-type (A) or sickle (S) alleles of the human β -globin gene. The left of each pair of tubes contains allele-specific primers to the wild-type alleles, the right tube primers to the sickle allele. The photograph was taken after 30 cycles of PCR, and the input DNAs and the alleles they contain are indicated. Fifty ng of DNA was used to begin PCR. Typing was done in triplicate (3 pairs of PCRs) for each input DNA.

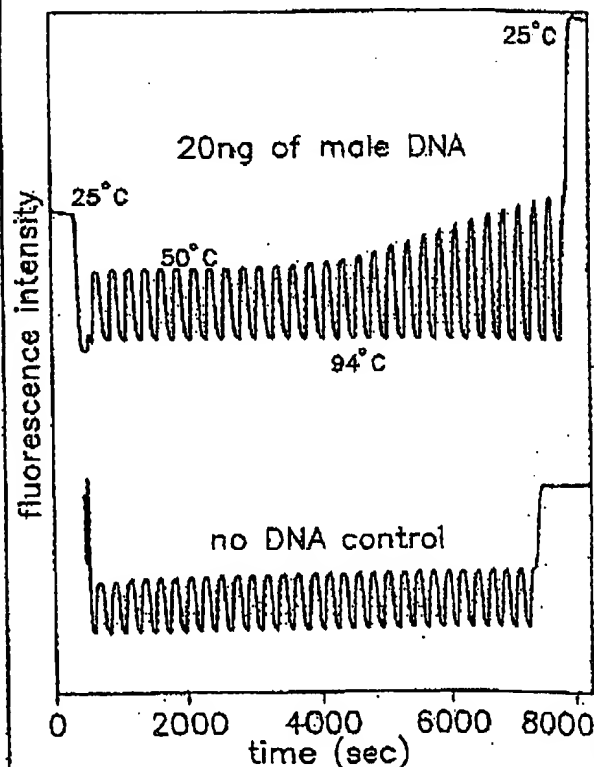


FIGURE 5 Continuous, real-time monitoring of a PCR. A fiber optic was used to carry excitation light to a PCR in progress and also emitted light back to a fluorometer (see Experimental Protocol). Amplification using human male-DNA specific primers in a PCR starting with 20 ng of human male DNA (top), or in a control PCR without DNA (bottom), were monitored. Thirty cycles of PCR were followed for each. The temperature cycled between 94°C (denaturation) and 50°C (annealing and extension). Note in the male DNA PCR, the cycle (time) dependent increase in fluorescence at the annealing/extension temperature.

DNA—up to microgram amounts—in order to have sufficient numbers of target sequences. This large amount of starting DNA in an amplification significantly increases the background fluorescence over which any additional fluorescence produced by PCR must be detected. An additional complication that occurs with targets in low copy-number is the formation of the "primer-dimer" artifact. This is the result of the extension of one primer using the other primer as a template. Although this occurs infrequently, once it occurs the extension product is a substrate for PCR amplification, and can compete with true PCR targets if those targets are rare. The primer-dimer product is of course dsDNA and thus is a potential source of false signal in this homogeneous assay.

To increase PCR specificity and reduce the effect of primer-dimer amplification, we are investigating a number of approaches, including the use of nested-primer amplifications that take place in a single tube³, and the "hot-start", in which nonspecific amplification is reduced by raising the temperature of the reaction before DNA synthesis begins²⁴. Preliminary results using these approaches suggest that primer-dimer is effectively reduced and it is possible to detect the increase in EtBr fluorescence in a PCR instigated by a single HIV genome in a background of 10^5 cells. With larger numbers of cells, the background fluorescence contributed by genomic DNA becomes problematic. To reduce this background, it may be possible to use sequence-specific DNA-binding dyes that can be made to preferentially bind PCR product over genomic DNA by incorporating the dye-binding DNA sequence into the PCR product through a 5' "add-on" to the oligonucleotide primer²⁴.

We have shown that the detection of fluorescence generated by an EtBr-containing PCR is straightforward, both once PCR is completed and continuously during thermocycling. The ease with which automation of specific DNA detection can be accomplished is the most promising aspect of this assay. The fluorescence analysis of completed PCRs is already possible with existing instrumentation in 96-well format²⁵. In this format, the fluorescence in each PCR can be quantitated before, after, and even at selected points during thermocycling by moving the rack of PCRs to a 96-microwell plate fluorescence reader²⁶.

The instrumentation necessary to continuously monitor multiple PCRs simultaneously is also simple in principle. A direct extension of the apparatus used here is to have multiple fiberoptics transmit the excitation light and fluorescent emissions to and from multiple PCRs. The ability to monitor multiple PCRs continuously may allow quantitation of target DNA copy number. Figure 3 shows that the larger the amount of starting target DNA, the sooner during PCR a fluorescence increase is detected. Preliminary experiments (Higuchi and Dollinger, manuscript in preparation) with continuous monitoring have shown a sensitivity to two-fold differences in initial target DNA concentration.

Conversely, if the number of target molecules is known—as it can be in genetic screening—continuous monitoring may provide a means of detecting false positive and false negative results. With a known number of target molecules, a true positive would exhibit detectable fluorescence by a predictable number of cycles of PCR. Increases in fluorescence detected before or after that cycle would indicate potential artifacts. False negative results due to, for example, inhibition of DNA polymerase, may be detected by including within each PCR an inefficiently amplifying marker. This marker results in a fluorescence increase only after a large number of cycles—many more than are necessary to detect a true

positive. If a sample fails to have a fluorescence increase after this many cycles, inhibition may be suspected. Since, in this assay, conclusions are drawn based on the presence or absence of fluorescence signal alone, such controls may be important. In any event, before any test based on this principle is ready for the clinic, an assessment of its false positive/false negative rates will need to be obtained using a large number of known samples.

In summary, the inclusion in PCR of dyes whose fluorescence is enhanced upon binding dsDNA makes it possible to detect specific DNA amplification from outside the PCR tube. In the future, instruments based upon this principle may facilitate the more widespread use of PCR in applications that demand the high throughput of samples.

EXPERIMENTAL PROTOCOL

Human HLA-DQ α gene amplifications containing EtBr. PCRs were set up in 100 μ l volumes containing 10 mM Tris-HCl, pH 8.3; 50 mM KCl; 4 mM MgCl₂; 2.5 units of *Taq* DNA polymerase (Perkin-Elmer Cetus, Norwalk, CT); 20 pmole each of human HLA-DQ α gene specific oligonucleotide primers (GH26 and GH27)¹⁹ and approximately 10^5 copies of DQ α PCR product diluted from a previous reaction. Ethidium bromide (EtBr; Sigma) was used at the concentrations indicated in Figure 2. Thermocycling proceeded for 20 cycles in a model 480 thermocycler (Perkin-Elmer Cetus, Norwalk, CT) using a "step-cycle" program of 94°C for 1 min., denaturation and 60°C for 30 sec., annealing and 72°C for 30 sec., extension.

Y-chromosome specific PCR. PCRs (100 μ l total reaction volume) containing 0.5 μ g/ml EtBr were prepared as described for HLA-DQ α , except with different primers and target DNAs. These PCRs contained 15 pmole each male DNA-specific primers Y1.1 and Y1.2²⁰, and either 60 ng male, 60 ng female, 2 ng male, or no human DNA. Thermocycling was 94°C for 1 min., and 60°C for 1 min using a "step-cycle" program. The number of cycles for a sample were as indicated in Figure 3. Fluorescence measurement is described below.

Allele-specific, human β -globin gene PCR. Amplifications of 100 μ l volume using 0.5 μ g/ml of EtBr were prepared as described for HLA-DQ α above except with different primers and target DNAs. These PCRs contained either primer pair HGP2/HB14A (wild-type globin specific primers) or HGP2/HB14S (sickle-globin specific primers) at 10 pmole each primer per PCR. These primers were developed by Wu et al.²¹. Three different target DNAs were used in separate amplifications—50 ng each of human DNA that was homozygous for the sickle trait (SS), DNA that was heterozygous for the sickle trait (AS), or DNA that was homozygous for the w.t. globin (AA). Thermocycling was for 30 cycles at 94°C for 1 min., and 55°C for 1 min., using a "step-cycle" program. An annealing temperature of 55°C had been shown by Wu et al.²¹ to provide allele-specific amplification. Completed PCRs were photographed through a red filter (Wratten 23A) after placing the reaction tubes atop a model TM-36 transilluminator (UV-products San Gabriel, CA).

Fluorescence measurement. Fluorescence measurements were made on PCRs containing EtBr in a Fluorolog-2 fluorometer (SPEX, Edison, NJ). Excitation was at the 500 nm band with about 2 nm bandwidth with a GG 435 nm cut-off filter (Melles Griest, Inc., Irvine, CA) to exclude second-order light. Emitted light was detected at 570 nm with a band width of about 7 nm. An OG 530 nm cut-off filter was used to remove the excitation light.

Continuous fluorescence monitoring of PCR. Continuous monitoring of a PCR in progress was accomplished using the spectrofluorometer and settings described above as well as a fiberoptic accessory (SPEX cat. no. 1950) to both send excitation light to, and receive emitted light from, a PCR placed in a well of a model 480 thermocycler (Perkin-Elmer Cetus). The probe end of the fiberoptic cable was attached with "5 minute-epoxy" to the open top of a PCR tube (a 0.5 ml polypropylene centrifuge tube with its cap removed) effectively sealing it. The exposed top of the PCR tube and the end of the fiberoptic cable were shielded from room light and the room lights were kept dimmed during each run. The monitored PCR was an amplification of Y-chromosome-specific repeat sequences as described above, except using an annealing/extension temperature of 50°C. The reaction was covered with mineral oil (2 drops) to prevent evaporation. Thermocycling and fluorescence measurement were started simultaneously. A time-base scan with a 10 second integration time

was used and the emission signal was ratioed to the excitation signal to control for changes in light-source intensity. Data were collected using the dm3000f, version 2.5 (SPEX) data system.

Acknowledgments

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RESEARCH

SIMULTANEOUS AMPLIFICATION AND DETECTION OF SPECIFIC DNA SEQUENCES

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We have enhanced the polymerase chain reaction (PCR) such that specific DNA sequences can be detected without opening the reaction tube. This enhancement requires the addition of ethidium bromide (EtBr) to a PCR. Since the fluorescence of EtBr increases in the presence of double-stranded (ds) DNA an increase in fluorescence in such a PCR indicates a positive amplification, which can be easily monitored externally. In fact, amplification can be continuously monitored in order to follow its progress. The ability to simultaneously amplify specific DNA sequences and detect the product of the amplification both simplifies and improves PCR and may facilitate its automation and more widespread use in the clinic or in other situations requiring high sample throughput.

Although the potential benefits of PCR¹ to clinical diagnostics are well known^{2,3}, it is still not widely used in this setting, even though it is four years since thermostable DNA polymerases⁴ made PCR practical. Some of the reasons for its slow acceptance are high cost, lack of automation of pre- and post-PCR processing steps, and false positive results from carryover-contamination. The first two points are related in that labor is the largest contributor to cost at the present stage of PCR development. Most current assays require some form of "downstream" processing once thermocycling is done in order to determine whether the target DNA sequence was present and has amplified. These include DNA hybridization^{5,6}, gel electrophoresis with or without use of restriction digestion^{7,8}, HPLC⁹, or capillary electrophoresis¹⁰. These methods are labor-intensive, have low throughput, and are difficult to automate. The third point is also closely related to downstream processing. The handling of the PCR product in these downstream processes increases the chances that amplified DNA will spread through the typing lab, resulting in a risk of

"carryover" false positives in subsequent testing¹¹.

These downstream processing steps would be eliminated if specific amplification and detection of amplified DNA took place simultaneously within an unopened reaction vessel. Assays in which such different processes take place without the need to separate reaction components have been termed "homogeneous". No truly homogeneous PCR assay has been demonstrated to date, although progress towards this end has been reported. Chehab, et al.¹², developed a PCR product detection scheme using fluorescent primers that resulted in a fluorescent PCR product. Allele-specific primers, each with different fluorescent tags, were used to indicate the genotype of the DNA. However, the unincorporated primers must still be removed in a downstream process in order to visualize the result. Recently, Holland, et al.¹³, developed an assay in which the endogenous 5' exonuclease assay of *Taq* DNA polymerase was exploited to cleave a labeled oligonucleotide probe. The probe would only cleave if PCR amplification had produced its complementary sequence. In order to detect the cleavage products, however, a subsequent process is again needed.

We have developed a truly homogeneous assay for PCR and PCR product detection based upon the greatly increased fluorescence that ethidium bromide and other DNA binding dyes exhibit when they are bound to ds-DNA¹⁴⁻¹⁶. As outlined in Figure 1, a prototypic PCR

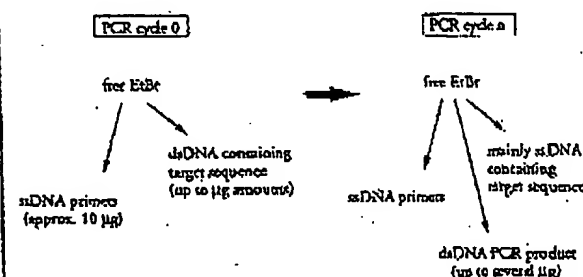


FIGURE 1 Principle of simultaneous amplification and detection of PCR product. The components of a PCR containing EtBr that are fluorescent are listed—EtBr itself, EtBr bound to either ssDNA or dsDNA. There is a large fluorescence enhancement when EtBr is bound to DNA and binding is greatly enhanced when DNA is double-stranded. After sufficient (n) cycles of PCR, the net increase in dsDNA results in additional EtBr binding, and a net increase in total fluorescence.

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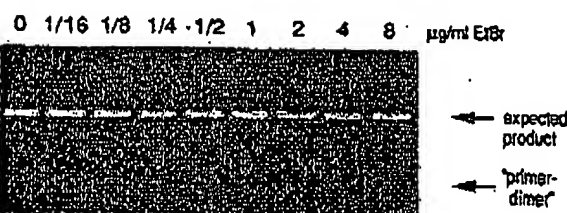


FIGURE 2 Gel electrophoresis of PCR amplification products of the human nuclear gene, HLA DQ α , made in the presence of increasing amounts of EtBr (up to 8 μ g/ml). The presence of EtBr has no obvious effect on the yield or specificity of amplification.

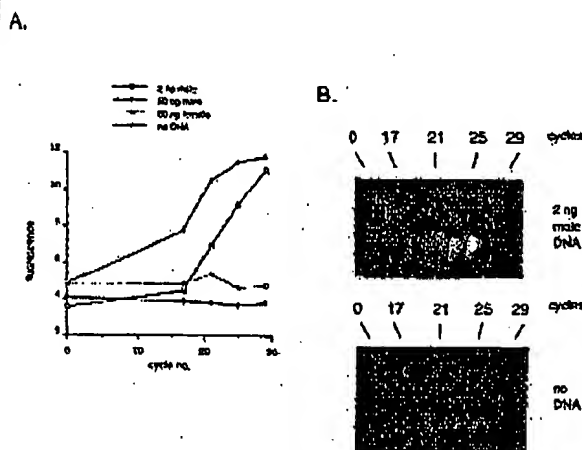


FIGURE 3 (A) Fluorescence measurements from PCR reactions that contain 0.5 μ g/ml EtBr and that are specific for Y-chromosome repeat sequences. Five replicate PCR reactions were begun containing each of the DNAs specified. At each indicated cycle, one of the five replicate PCR reactions for each DNA was removed from thermocycling and its fluorescence measured. Units of fluorescence are arbitrary. (B) UV photograph of PCR tubes (0.5 ml Eppendorf-style, polypropylene micro-centrifuge tubes) containing reactions, those starting from 2 ng male DNA and control reactions without any DNA, from (A).

begins with primers that are single-stranded DNA (ssDNA), dNTPs, and DNA polymerase. An amount of dsDNA containing the target sequence (target DNA) is also typically present. This amount can vary, depending on the application, from single-cell amounts of DNA¹⁷ to micrograms per PCR¹⁸. If EtBr is present, the reagents that will fluoresce, in order of increasing fluorescence, are free EtBr itself, and EtBr bound to the single-stranded DNA primers and to the double-stranded target DNA (by its intercalation between the stacked bases of the DNA double-helix). After the first denaturation cycle, target DNA will be largely single-stranded. After a PCR is completed, the most significant change is the increase in the amount of dsDNA (the PCR product itself) of up to several micrograms. Formerly free EtBr is bound to the additional dsDNA, resulting in an increase in fluorescence. There is also some decrease in the amount of ssDNA primer, but because the binding of EtBr to ssDNA is much less than to dsDNA, the effect of this change on the total fluorescence of the sample is small. The fluorescence increase can be measured by directing excitation illumination through the walls of the amplification vessel

before and after, or even continuously during, thermocycling.

RESULTS

PCR in the presence of EtBr. In order to assess the effect of EtBr in PCR, amplifications of the human HLA DQ α gene¹⁹ were performed with the dye present at concentrations from 0.06 to 8.0 μ g/ml (a typical concentration of EtBr used in staining of nucleic acids following gel electrophoresis is 0.5 μ g/ml). As shown in Figure 2, gel electrophoresis revealed little or no difference in the yield or quality of the amplification product whether EtBr was absent or present at any of these concentrations, indicating that EtBr does not inhibit PCR.

Detection of human Y-chromosome specific sequences. Sequence-specific, fluorescence enhancement of EtBr as a result of PCR was demonstrated in a series of amplifications containing 0.5 μ g/ml EtBr and primers specific to repeat DNA sequences found on the human Y-chromosome²⁰. These PCRs initially contained either 60 ng male, 60 ng female, 2 ng male human or no DNA. Five replicate PCRs were begun for each DNA. After 0, 17, 21, 24 and 29 cycles of thermocycling, a PCR for each DNA was removed from the thermocycler, and its fluorescence measured in a spectrofluorometer and plotted vs. amplification cycle number (Fig. 3A). The shape of this curve reflects the fact that by the time an increase in fluorescence can be detected, the increase in DNA is becoming linear and not exponential with cycle number. As shown, the fluorescence increased about three-fold over the background fluorescence for the PCRs containing human male DNA, but did not significantly increase for negative control PCRs, which contained either no DNA or human female DNA. The more male DNA present to begin with—60 ng versus 2 ng—the fewer cycles were needed to give a detectable increase in fluorescence. Gel electrophoresis on the products of these amplifications showed that DNA fragments of the expected size were made in the male DNA containing reactions and that little DNA synthesis took place in the control samples.

In addition, the increase in fluorescence was visualized by simply laying the completed, unopened PCRs on a UV transilluminator and photographing them through a red filter. This is shown in figure 3B for the reactions that began with 2 ng male DNA and those with no DNA.

Detection of specific alleles of the human β -globin gene. In order to demonstrate that this approach has adequate specificity to allow genetic screening, a detection of the sickle-cell anemia mutation was performed. Figure 4 shows the fluorescence from completed amplifications containing EtBr (0.5 μ g/ml) as detected by photography of the reaction tubes on a UV transilluminator. These reactions were performed using primers specific for either the wild-type or sickle-cell mutation of the human β -globin gene²¹. The specificity for each allele is imparted by placing the sickle-mutation site at the terminal 3' nucleotide of one primer. By using an appropriate primer annealing temperature, primer extension—and thus amplification—can take place only if the 3' nucleotide of the primer is complementary to the β -globin allele present.²²

Each pair of amplifications shown in Figure 4 consists of a reaction with either the wild-type allele specific (left tube) or sickle-allele specific (right tube) primers. Three different DNAs were typed: DNA from a homozygous, wild-type β -globin individual (AA); from a heterozygous sickle β -globin individual (AS); and from a homozygous sickle β -globin individual (SS). Each DNA (50 ng genomic DNA to start each PCR) was analyzed in triplicate (3 pairs

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of reactions each). The DNA type was reflected in the relative fluorescence intensities in each pair of completed amplifications. There was a significant increase in fluorescence only where a β -globin allele DNA matched the primer set. When measured on a spectrofluorometer (data not shown), this fluorescence was about three times that present in a PCR where both β -globin alleles were mismatched to the primer set. Gel electrophoresis (not shown) established that this increase in fluorescence was due to the synthesis of nearly a microgram of a DNA fragment of the expected size for β -globin. There was little synthesis of dsDNA in reactions in which the allele-specific primer was mismatched to both alleles.

Continuous monitoring of a PCR. Using a fiber optic device, it is possible to direct excitation illumination from a spectrofluorometer to a PCR undergoing thermocycling and to return its fluorescence to the spectrofluorometer. The fluorescence readout of such an arrangement, directed at an EtBr-containing amplification of Y-chromosome specific sequences from 25 ng of human male DNA, is shown in Figure 5. The readout from a control PCR with no target DNA is also shown. Thirty cycles of PCR were monitored for each.

The fluorescence trace as a function of time clearly shows the effect of the thermocycling. Fluorescence intensity rises and falls inversely with temperature. The fluorescence intensity is minimum at the denaturation temperature (94°C) and maximum at the annealing/extension temperature (50°C). In the negative-control PCR, these fluorescence maxima and minima do not change significantly over the thirty thermocycles, indicating that there is little dsDNA synthesis without the appropriate target DNA, and there is little if any bleaching of EtBr during the continuous illumination of the sample.

In the PCR containing male DNA, the fluorescence maxima at the annealing/extension temperature begin to increase at about 4000 seconds of thermocycling, and continue to increase with time, indicating that dsDNA is being produced at a detectable level. Note that the fluorescence minima at the denaturation temperature do not significantly increase, presumably because at this temperature there is no dsDNA for EtBr to bind. Thus the course of the amplification is followed by tracking the fluorescence increase at the annealing temperature. Analysis of the products of these two amplifications by gel electrophoresis showed a DNA fragment of the expected size for the male DNA containing sample and no detectable DNA synthesis for the control sample.

DISCUSSION

Downstream processes such as hybridization to a sequence-specific probe can enhance the specificity of DNA detection by PCR. The elimination of these processes means that the specificity of this homogeneous assay depends solely on that of PCR. In the case of sickle-cell disease, we have shown that PCR alone has sufficient DNA sequence specificity to permit genetic screening. Using appropriate amplification conditions, there is little non-specific production of dsDNA in the absence of the appropriate target allele.

The specificity required to detect pathogens can be more or less than that required to do genetic screening, depending on the number of pathogens in the sample and the amount of other DNA that must be taken with the sample. A difficult target is HIV, which requires detection of a viral genome that can be at the level of a few copies per thousands of host cells⁶. Compared with genetic screening, which is performed on cells containing at least one copy of the target sequence, HIV detection requires both more specificity and the input of more total

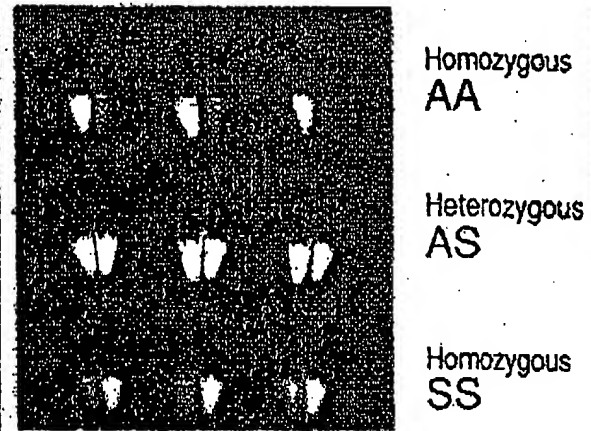


FIGURE 4 UV photograph of PCR tubes containing amplifications using EtBr that are specific to wild-type (A) or sickle (S) alleles of the human β -globin gene. The left of each pair of tubes contains allele-specific primers to the wild-type alleles, the right tube primers to the sickle allele. The photograph was taken after 30 cycles of PCR, and the input DNAs and the alleles they contain are indicated. Fifty ng of DNA was used to begin PCR. Typing was done in triplicate (3 pairs of PCRs) for each input DNA.

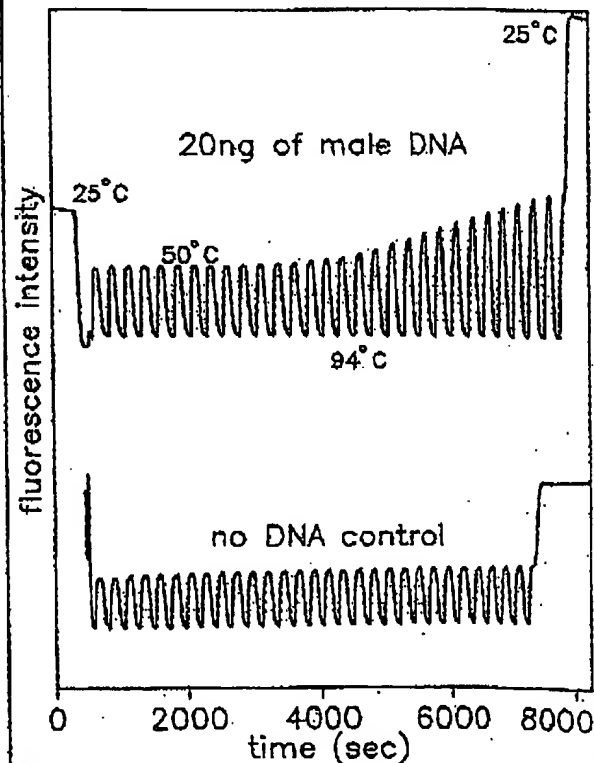


FIGURE 5 Continuous, real-time monitoring of a PCR. A fiber optic was used to carry excitation light to a PCR in progress and also emitted light back to a fluorometer (see Experimental Protocol). Amplification using human male-DNA specific primers in a PCR starting with 20 ng of human male DNA (top), or in a control PCR without DNA (bottom), were monitored. Thirty cycles of PCR were followed for each. The temperature cycled between 94°C (denaturation) and 50°C (annealing and extension). Note in the male DNA PCR, the cycle (time) dependent increase in fluorescence at the annealing/extension temperature.

DNA—up to microgram amounts—in order to have sufficient numbers of target sequences. This large amount of starting DNA in an amplification significantly increases the background fluorescence over which any additional fluorescence produced by PCR must be detected. An additional complication that occurs with targets in low copy-number is the formation of the "primer-dimer" artifact. This is the result of the extension of one primer using the other primer as a template. Although this occurs infrequently, once it occurs the extension product is a substrate for PCR amplification, and can compete with true PCR targets if those targets are rare. The primer-dimer product is of course dsDNA and thus is a potential source of false signal in this homogeneous assay.

To increase PCR specificity and reduce the effect of primer-dimer amplification, we are investigating a number of approaches, including the use of nested-primer amplifications that take place in a single tube³, and the "hot-start", in which nonspecific amplification is reduced by raising the temperature of the reaction before DNA synthesis begins²⁴. Preliminary results using these approaches suggest that primer-dimer is effectively reduced and it is possible to detect the increase in EtBr fluorescence in a PCR instigated by a single HIV genome in a background of 10^5 cells. With larger numbers of cells, the background fluorescence contributed by genomic DNA becomes problematic. To reduce this background, it may be possible to use sequence-specific DNA-binding dyes that can be made to preferentially bind PCR product over genomic DNA by incorporating the dye-binding DNA sequence into the PCR product through a 5' "add-on" to the oligonucleotide primer²⁴.

We have shown that the detection of fluorescence generated by an EtBr-containing PCR is straightforward, both once PCR is completed and continuously during thermocycling. The ease with which automation of specific DNA detection can be accomplished is the most promising aspect of this assay. The fluorescence analysis of completed PCRs is already possible with existing instrumentation in 96-well format²⁵. In this format, the fluorescence in each PCR can be quantitated before, after, and even at selected points during thermocycling by moving the rack of PCRs to a 96-microwell plate fluorescence reader²⁶.

The instrumentation necessary to continuously monitor multiple PCRs simultaneously is also simple in principle. A direct extension of the apparatus used here is to have multiple fiberoptics transmit the excitation light and fluorescent emissions to and from multiple PCRs. The ability to monitor multiple PCRs continuously may allow quantitation of target DNA copy number. Figure 3 shows that the larger the amount of starting target DNA, the sooner during PCR a fluorescence increase is detected. Preliminary experiments (Higuchi and Dollinger, manuscript in preparation) with continuous monitoring have shown a sensitivity to two-fold differences in initial target DNA concentration.

Conversely, if the number of target molecules is known—as it can be in genetic screening—continuous monitoring may provide a means of detecting false positive and false negative results. With a known number of target molecules, a true positive would exhibit detectable fluorescence by a predictable number of cycles of PCR. Increases in fluorescence detected before or after that cycle would indicate potential artifacts. False negative results due to, for example, inhibition of DNA polymerase, may be detected by including within each PCR an inefficiently amplifying marker. This marker results in a fluorescence increase only after a large number of cycles—many more than are necessary to detect a true

positive. If a sample fails to have a fluorescence increase after this many cycles, inhibition may be suspected. Since, in this assay, conclusions are drawn based on the presence or absence of fluorescence signal alone, such controls may be important. In any event, before any test based on this principle is ready for the clinic, an assessment of its false positive/false negative rates will need to be obtained using a large number of known samples.

In summary, the inclusion in PCR of dyes whose fluorescence is enhanced upon binding dsDNA makes it possible to detect specific DNA amplification from outside the PCR tube. In the future, instruments based upon this principle may facilitate the more widespread use of PCR in applications that demand the high throughput of samples.

EXPERIMENTAL PROTOCOL

Human HLA-DQ α gene amplifications containing EtBr. PCRs were set up in 100 μ l volumes containing 10 mM Tris-HCl, pH 8.3; 50 mM KCl; 4 mM MgCl₂; 2.5 units of *Taq* DNA polymerase (Perkin-Elmer Cetus, Norwalk, CT); 20 pmole each of human HLA-DQ α gene specific oligonucleotide primers GH26 and GH27¹⁹ and approximately 10^5 copies of DQ α PCR product diluted from a previous reaction. Ethidium bromide (EtBr; Sigma) was used at the concentrations indicated in Figure 2. Thermocycling proceeded for 20 cycles in a model 480 thermocycler (Perkin-Elmer Cetus, Norwalk, CT) using a "step-cycle" program of 94°C for 1 min., denaturation and 60°C for 30 sec., annealing and 72°C for 30 sec., extension.

Y-chromosome specific PCR. PCRs (100 μ l total reaction volume) containing 0.5 μ g/ml EtBr were prepared as described for HLA-DQ α , except with different primers and target DNAs. These PCRs contained 15 pmole each male DNA-specific primers Y1.1 and Y1.2²⁰, and either 60 ng male, 60 ng female, 2 ng male, or no human DNA. Thermocycling was 94°C for 1 min. and 60°C for 1 min using a "step-cycle" program. The number of cycles for a sample were as indicated in Figure 3. Fluorescence measurement is described below.

Allele-specific, human β -globin gene PCR. Amplifications of 100 μ l volume using 0.5 μ g/ml of EtBr were prepared as described for HLA-DQ α above except with different primers and target DNAs. These PCRs contained either primer pair HGP2/HB14A (wild-type globin specific primers) or HGP2/HB14S (sickle-globin specific primers) at 10 pmole each primer per PCR. These primers were developed by Wu et al.²¹. Three different target DNAs were used in separate amplifications—50 ng each of human DNA that was homozygous for the sickle trait (SS), DNA that was heterozygous for the sickle trait (AS), or DNA that was homozygous for the w.t. globin (AA). Thermocycling was for 30 cycles at 94°C for 1 min. and 55°C for 1 min. using a "step-cycle" program. An annealing temperature of 55°C had been shown by Wu et al.²¹ to provide allele-specific amplification. Completed PCRs were photographed through a red filter (Wratten 23A) after placing the reaction tubes atop a model TM-36 (transilluminator (UV-products San-Gabriel, CA).

Fluorescence measurement. Fluorescence measurements were made on PCRs containing EtBr in a Fluorolog-2 fluorometer (SPEX, Edison, NJ). Excitation was at the 500 nm band with about 2 nm bandwidth with a GG 435 nm cut-off filter (Melles Griest, Inc., Irvine, CA) to exclude second-order light. Emitted light was detected at 570 nm with a bandwidth of about 7 nm. An OG 530 nm cut-off filter was used to remove the excitation light.

Continuous fluorescence monitoring of PCR. Continuous monitoring of a PCR in progress was accomplished using the spectrofluorometer and settings described above as well as a fiberoptic accessory (SPEX cat. no. 1950) to both send excitation light to, and receive emitted light from, a PCR placed in a well of a model 480 thermocycler (Perkin-Elmer Cetus). The probe end of the fiberoptic cable was attached with "5 minute-epoxy" to the open top of a PCR tube (a 0.5 ml polypropylene centrifuge tube with its cap removed) effectively sealing it. The exposed top of the PCR tube and the end of the fiberoptic cable were shielded from room light and the room lights were kept dimmed during each run. The monitored PCR was an amplification of Y-chromosome-specific repeat sequences as described above, except using an annealing/extension temperature of 50°C. The reaction was covered with mineral oil (2 drops) to prevent evaporation. Thermocycling and fluorescence measurement were started simultaneously. A time-base scan with a 10 second integration time

was used and the emission signal was ratioed to the excitation signal to control for changes in light-source intensity. Data were collected using the dm3000f, version 2.5 (SPEX) data system.

Acknowledgments

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Oligonucleotides with Fluorescent Dyes at Opposite Ends Provide a Quenched Probe System Useful for Detecting PCR Product and Nucleic Acid Hybridization

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The 5' nuclease PCR assay detects the accumulation of specific PCR product by hybridization and cleavage of a double-labeled fluorogenic probe during the amplification reaction. The probe is an oligonucleotide with both a reporter fluorescent dye and a quencher dye attached. An increase in reporter fluorescence intensity indicates that the probe has hybridized to the target PCR product and has been cleaved by the 5' → 3' nucleolytic activity of *Taq* DNA polymerase. In this study, probes with the quencher dye attached to an internal nucleotide were compared with probes with the quencher dye attached to the 3'-end nucleotide. In all cases, the reporter dye was attached to the 5' end. All intact probes showed quenching of the reporter fluorescence. In general, probes with the quencher dye attached to the 3'-end nucleotide exhibited a larger signal in the 5' nuclease PCR assay than the internally labeled probes. It is proposed that the larger signal is caused by increased likelihood of cleavage by *Taq* DNA polymerase when the probe is hybridized to a template strand during PCR. Probes with the quencher dye attached to the 3'-end nucleotide also exhibited an increase in reporter fluorescence intensity when hybridized to a complementary strand. Thus, oligonucleotides with reporter and quencher dyes attached at opposite ends can be used as homogeneous hybridization probes.

A homogeneous assay for detecting the accumulation of specific PCR product that uses a double-labeled fluorogenic probe was described by Lee et al.⁽¹⁾ The assay exploits the 5' → 3' nucleolytic activity of *Taq* DNA polymerase^(2,3) and is diagrammed in Figure 1. The fluorogenic probe consists of an oligonucleotide with a reporter fluorescent dye, such as a fluorescein, attached to the 5' end; and a quencher dye, such as a rhodamine, attached internally. When the fluorescein is excited by irradiation, its fluorescent emission will be quenched if the rhodamine is close enough to be excited through the process of fluorescence energy transfer (FET).^(4,5) During PCR, if the probe is hybridized to a template strand, *Taq* DNA polymerase will cleave the probe because of its inherent 5' → 3' nucleolytic activity. If the cleavage occurs between the fluorescein and rhodamine dyes, it causes an increase in fluorescein fluorescence intensity because the fluorescein is no longer quenched. The increase in fluorescein fluorescence intensity indicates that the probe-specific PCR product has been generated. Thus, FET between a reporter dye and a quencher dye is critical to the performance of the probe in the 5' nuclease PCR assay.

Quenching is completely dependent on the physical proximity of the two dyes.⁽⁶⁾ Because of this, it has been assumed that the quencher dye must be attached near the 5' end. Surprisingly, we have found that attaching a rhodamine dye at the 3' end of a probe still provides adequate quenching for the probe to perform in the 5' nuclease

PCR assay. Furthermore, cleavage of this type of probe is not required to achieve some reduction in quenching. Oligonucleotides with a reporter dye on the 5' end and a quencher dye on the 3' end exhibit a much higher reporter fluorescence when double-stranded as compared with single-stranded. This should make it possible to use this type of double-labeled probe for homogeneous detection of nucleic acid hybridization.

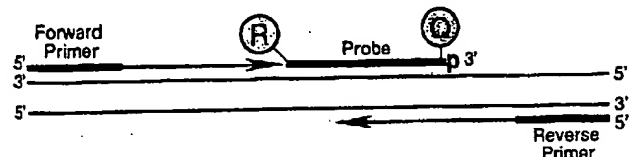
MATERIALS AND METHODS

Oligonucleotides

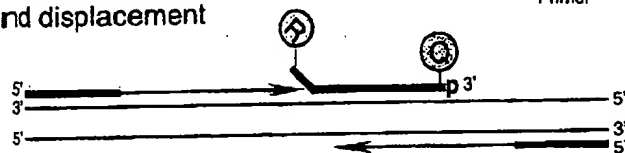
Table 1 shows the nucleotide sequence of the oligonucleotides used in this study. Linker arm nucleotide (LAN) phosphoramidite was obtained from Glen Research. The standard DNA phosphoramidites, 6-carboxyfluorescein (6-FAM) phosphoramidite, 6-carboxytetramethylrhodamine succinimidyl ester (TAMRA NHS ester), and Phosphalink for attaching a 3'-blocking phosphate, were obtained from Perkin-Elmer, Applied Biosystems Division. Oligonucleotide synthesis was performed using an ABI model 394 DNA synthesizer (Applied Biosystems). Primer and complement oligonucleotides were purified using Oligo Purification Cartridges (Applied Biosystems). Double-labeled probes were synthesized with 6-FAM-labeled phosphoramidite at the 5' end, LAN replacing one of the T's in the sequence, and Phosphalink at the 3' end. Following deprotection and ethanol precipitation, TAMRA NHS ester was coupled to the LAN-containing oligonucleotide in 250

Research

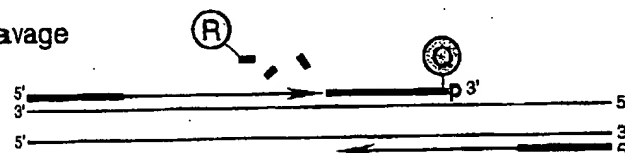
Polymerization



Strand displacement



Cleavage



Polymerization completed

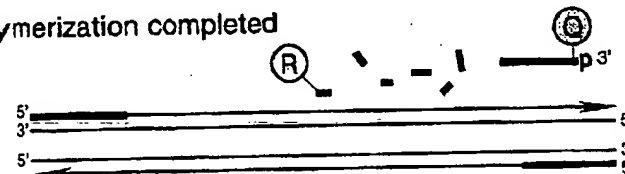


FIGURE 1 Diagram of 5' nuclease assay. Stepwise representation of the 5' → 3' nucleolytic activity of *Taq* DNA polymerase acting on a fluorogenic probe during one extension phase of PCR.

mm Na-bicarbonate buffer (pH 9.0) at room temperature. Unreacted dye was removed by passage over a PD-10 Sephadex column. Finally, the double-labeled probe was purified by preparative high-performance liquid chromatography (HPLC) using an Aquapore C₈ 220×4.6-mm column with 7-μm particle size. The column was developed with a 24-min linear gradient of 8–20% acetonitrile in 0.1 M TEAA (triethylamine acetate). Probes are named by designating the sequence from Table 1 and the position of the LAN-TAMRA moiety. For example, probe A1-7 has sequence A1 with LAN-TAMRA at nucleotide position 7 from the 5' end.

PCR Systems

All PCR amplifications were performed in the Perkin-Elmer GeneAmp PCR System 9600 using 50-μl reactions that contained 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 200 μM dATP, 200 μM dCTP, 200 μM dGTP, 400 μM dUTP, 0.5 unit of AmpErase uracil N-glycosylase (Perkin-Elmer), and 1.25 unit of AmpliTaq DNA polymerase (Perkin-Elmer). A 295-bp segment from exon 3 of the human β-actin

gene (nucleotides 2141–2435 in the sequence of Nakajima-Iijima et al.)⁽⁷⁾ was amplified using primers AFP and ARP (Table 1), which are modified slightly from those of du Breuil et al.⁽⁸⁾ Actin amplification reactions contained 4 mM MgCl₂, 20 ng of human genomic DNA, 50 nM A1 or A3 probe, and 300 nM each

primer. The thermal regimen was 50°C (2 min), 95°C (10 min), 40 cycles of 95°C (20 sec), 60°C (1 min), and hold at 72°C. A 515-bp segment was amplified from a plasmid that consists of a segment of λ DNA (nucleotides 32,220–32,747) inserted in the *Sma*I site of vector pUC119. These reactions contained 3.5 mM MgCl₂, 1 ng of plasmid DNA, 50 nM P2 or P5 probe, 200 nM primer F119, and 200 nM primer R119. The thermal regimen was 50°C (2 min), 95°C (10 min), 25 cycles of 95°C (20 sec), 57°C (1 min), and hold at 72°C.

Fluorescence Detection

For each amplification reaction, a 40-μl aliquot of a sample was transferred to an individual well of a white, 96-well microtiter plate (Perkin-Elmer). Fluorescence was measured on the Perkin-Elmer TaqMan LS-50B System, which consists of a luminescence spectrometer with plate reader assembly, a 485-nm excitation filter, and a 515-nm emission filter. Excitation was at 488 nm using a 5-nm slit width. Emission was measured at 518 nm for 6-FAM (the reporter or R value) and 582 nm for TAMRA (the quencher or Q value) using a 10-nm slit width. To determine the increase in reporter emission that is caused by cleavage of the probe during PCR, three normalizations are applied to the raw emission data. First, emission intensity of a buffer blank is subtracted for each wavelength. Second, emission intensity of the reporter is

TABLE 1 Sequences of Oligonucleotides

Name	Type	Sequence
F119	primer	ACCCACAGGAAGTATGATCACCACCTC
R119	primer	ATGTCGCGTTCCGGCTGACGTTCTGC
P2	probe	TCGCATTACTGATCGTTGCCAACCAGTp
P2C	complement	GTACTGGTTGGCAACGATCAGTAATGCGATG
P5	probe	CGGA'TTTGCTGGTATCTATGACAAGGATp
P5C	complement	TTCATCCTTGTCATAGATACCAGCAAATCCG
AFP	primer	TCACCCACACTGTGCCCATCTACGA
ARP	primer	CAGCGGAACCGCTCATTGCCAATGG
A1	probe	ATGCCCTCCCCATGCCATCTCTGCGTp
A1C	complement	AGACGCAGGATGGCATGGGGAGGGCATAC
A3	probe	CGCCCTGGACTTCGAGCAAGAGATp
A3C	complement	CCATCTCTTGCTCGAAGTCCAGGGCGAC

For each oligonucleotide used in this study, the nucleic acid sequence is given, written in the 5' → 3' direction. There are three types of oligonucleotides: PCR primer, fluorogenic probe used in the 5' nuclease assay, and complement used to hybridize to the corresponding probe. For the probes, the underlined base indicates a position where LAN with TAMRA attached was substituted for a T. (p) The presence of a 3' phosphate on each probe.

A1-2 RAQCCCTCCCCATGCCATCCTGCGTp
 A1-7 RATGCCCQCCCCATGCCATCCTGCGTp
 A1-14 RATGCCCTCCCCCAQGCCATCCTGCGTp
 A1-19 RATGCCCTCCCCATGCCAQCTGCGTp
 A1-22 RATGCCCTCCCCATGCCATCCQCGTp
 A1-26 RATGCCCTCCCCATGCCATCCTGCGQp

Probe	518 nm		582 nm		RQ ⁻	RQ ⁺	Δ RQ
	no temp.	+ temp.	no temp.	+ temp.			
A1-2	25.5 \pm 2.1	32.7 \pm 1.9	38.2 \pm 3.0	38.2 \pm 2.0	0.67 \pm 0.01	0.86 \pm 0.06	0.19 \pm 0.06
A1-7	53.5 \pm 4.3	395.1 \pm 21.4	108.5 \pm 6.3	110.3 \pm 5.3	0.49 \pm 0.03	3.58 \pm 0.17	3.09 \pm 0.18
A1-14	127.0 \pm 4.9	403.5 \pm 19.1	109.7 \pm 5.3	93.1 \pm 6.3	1.16 \pm 0.02	4.34 \pm 0.15	3.18 \pm 0.15
A1-19	187.5 \pm 17.9	422.7 \pm 7.7	70.3 \pm 7.4	73.0 \pm 2.8	2.67 \pm 0.05	5.80 \pm 0.15	3.13 \pm 0.16
A1-22	224.6 \pm 9.4	482.2 \pm 43.6	100.0 \pm 4.0	96.2 \pm 9.6	2.25 \pm 0.03	5.02 \pm 0.11	2.77 \pm 0.12
A1-26	160.2 \pm 8.9	454.1 \pm 18.4	93.1 \pm 5.4	90.7 \pm 3.2	1.72 \pm 0.02	5.01 \pm 0.08	3.29 \pm 0.08

FIGURE 2 Results of 5' nuclease assay comparing β -actin probes with TAMRA at different nucleotide positions. As described in Materials and Methods, PCR amplifications containing the indicated probes were performed, and the fluorescence emission was measured at 518 and 582 nm. Reported values are the average \pm 1 S.D. for six reactions run without added template (no temp.) and six reactions run with template (+temp.). The RQ ratio was calculated for each individual reaction and averaged to give the reported RQ⁻ and RQ⁺ values.

divided by the emission intensity of the quencher to give an RQ ratio for each reaction tube. This normalizes for well-to-well variations in probe concentration and fluorescence measurement. Finally, Δ RQ is calculated by subtracting the RQ value of the no-template control (RQ⁻) from the RQ value for the complete reaction including template (RQ⁺).

RESULTS

A series of probes with increasing distances between the fluorescein reporter and rhodamine quencher were tested to investigate the minimum and maximum spacing that would give an acceptable performance in the 5' nuclease PCR assay. These probes hybridize to a target

sequence in the human β -actin gene. Figure 2 shows the results of an experiment in which these probes were included in PCR that amplified a segment of the β -actin gene containing the target sequence. Performance in the 5' nuclease PCR assay is monitored by the magnitude of Δ RQ, which is a measure of the increase in reporter fluorescence caused by PCR amplification of the probe target. Probe A1-2 has a Δ RQ value that is close to zero, indicating that the probe was not cleaved appreciably during the amplification reaction. This suggests that with the quencher dye on the second nucleotide from the 5' end, there is insufficient room for *Taq* polymerase to cleave efficiently between the reporter and quencher. The other five probes exhibited comparable Δ RQ values that are

clearly different from zero. Thus, all five probes are being cleaved during PCR amplification resulting in a similar increase in reporter fluorescence. It should be noted that complete digestion of a probe produces a much larger increase in reporter fluorescence than that observed in Figure 2 (data not shown). Thus, even in reactions where amplification occurs, the majority of probe molecules remain uncleaved. It is mainly for this reason that the fluorescence intensity of the quencher dye TAMRA changes little with amplification of the target. This is what allows us to use the 582-nm fluorescence reading as a normalization factor.

The magnitude of RQ⁻ depends mainly on the quenching efficiency inherent in the specific structure of the probe and the purity of the oligonucleotide. Thus, the larger RQ⁻ values indicate that probes A1-14, A1-19, A1-22, and A1-26 probably have reduced quenching as compared with A1-7. Still, the degree of quenching is sufficient to detect a highly significant increase in reporter fluorescence when each of these probes is cleaved during PCR.

To further investigate the ability of TAMRA on the 3' end to quench 6-FAM on the 5' end, three additional pairs of probes were tested in the 5' nuclease PCR assay. For each pair, one probe has TAMRA attached to an internal nucleotide and the other has TAMRA attached to the 3' end nucleotide. The results are shown in Table 2. For all three sets, the probe with the 3' quencher exhibits a Δ RQ value that is considerably higher than for the probe with the internal quencher. The RQ⁻ values suggest that differences in quenching are not as great as those observed with some of the A1 probes. These results demonstrate that a quencher dye on the 3' end of an oligonucleotide can quench efficiently the

TABLE 2 Results of 5' Nuclease Assay Comparing Probes with TAMRA Attached to an Internal or 3'-terminal Nucleotide

Probe	518 nm		582 nm		RQ ⁻	RQ ⁺	Δ RQ
	no temp.	+ temp.	no temp.	+ temp.			
A3-6	54.6 \pm 3.2	84.8 \pm 3.7	116.2 \pm 6.4	115.6 \pm 2.5	0.47 \pm 0.02	0.73 \pm 0.03	0.26 \pm 0.04
A3-24	72.1 \pm 2.9	236.5 \pm 11.1	84.2 \pm 4.0	90.2 \pm 3.8	0.86 \pm 0.02	2.62 \pm 0.05	1.76 \pm 0.05
P2-7	82.8 \pm 4.4	384.0 \pm 34.1	105.1 \pm 6.4	120.4 \pm 10.2	0.79 \pm 0.02	3.19 \pm 0.16	2.40 \pm 0.16
P2-27	113.4 \pm 6.6	555.4 \pm 14.1	140.7 \pm 8.5	118.7 \pm 4.8	0.81 \pm 0.01	4.68 \pm 0.10	3.88 \pm 0.10
P5-10	77.5 \pm 6.5	244.4 \pm 15.9	86.7 \pm 4.3	95.8 \pm 6.7	0.89 \pm 0.05	2.55 \pm 0.06	1.66 \pm 0.08
P5-28	64.0 \pm 5.2	333.6 \pm 12.1	100.6 \pm 6.1	94.7 \pm 6.3	0.63 \pm 0.02	3.53 \pm 0.12	2.89 \pm 0.13

Reactions containing the indicated probes and calculations were performed as described in Material and Methods and in the legend to Fig. 2.

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fluorescence of a reporter dye on the 5' end. The degree of quenching is sufficient for this type of oligonucleotide to be used as a probe in the 5' nuclease PCR assay.

To test the hypothesis that quenching by a 3' TAMRA depends on the flexibility of the oligonucleotide, fluorescence was measured for probes in the single-stranded and double-stranded states. Table 3 reports the fluorescence observed at 518 and 582 nm. The relative degree of quenching is assessed by calculating the RQ ratio. For probes with TAMRA 6–10 nucleotides from the 5' end, there is little difference in the RQ values when comparing single-stranded with double-stranded oligonucleotides. The results for probes with TAMRA at the 3' end are much different. For these probes, hybridization to a complementary strand causes a dramatic increase in RQ. We propose that this loss of quenching is caused by the rigid structure of double-stranded DNA, which prevents the 5' and 3' ends from being in proximity.

When TAMRA is placed toward the 3' end, there is a marked Mg^{2+} effect on quenching. Figure 3 shows a plot of observed RQ values for the A1 series of probes as a function of Mg^{2+} concentration. With TAMRA attached near the 5' end (probe A1-2 or A1-7), the RQ value at 0 mM Mg^{2+} is only slightly higher than RQ at 10 mM Mg^{2+} . For probes A1-19, A1-22, and A1-26, the RQ values at 0 mM Mg^{2+} are very high, indicating a much

reduced quenching efficiency. For each of these probes, there is a marked decrease in RQ at 1 mM Mg^{2+} followed by a gradual decline as the Mg^{2+} concentration increases to 10 mM. Probe A1-14 shows an intermediate RQ value at 0 mM Mg^{2+} with a gradual decline at higher Mg^{2+} concentrations. In a low-salt environment with no Mg^{2+} present, a single-stranded oligonucleotide would be expected to adopt an extended conformation because of electrostatic repulsion. The binding of Mg^{2+} ions acts to shield the negative charge of the phosphate backbone so that the oligonucleotide can adopt conformations where the 3' end is close to the 5' end. Therefore, the observed Mg^{2+} effects support the notion that quenching of a 5' reporter dye by TAMRA at or near the 3' end depends on the flexibility of the oligonucleotide.

DISCUSSION

The striking finding of this study is that it seems the rhodamine dye TAMRA, placed at any position in an oligonucleotide, can quench the fluorescent emission of a fluorescein (6-FAM) placed at the 5' end. This implies that a single-stranded, double-labeled oligonucleotide must be able to adopt conformations where the TAMRA is close to the 5' end. It should be noted that the decay of 6-FAM in the excited state requires a certain amount of time. Therefore, what

matters for quenching is not the average distance between 6-FAM and TAMRA but, rather, how close TAMRA can get to 6-FAM during the lifetime of the 6-FAM excited state. As long as the decay time of the excited state is relatively long compared with the molecular motions of the oligonucleotide, quenching can occur. Thus, we propose that TAMRA at the 3' end, or any other position, can quench 6-FAM at the 5' end because TAMRA is in proximity to 6-FAM often enough to be able to accept energy transfer from an excited 6-FAM.

Details of the fluorescence measurements remain puzzling. For example, Table 3 shows that hybridization of probes A1-26, A3-24, and P5-28 to their complementary strands not only causes a large increase in 6-FAM fluorescence at 518 nm but also causes a modest increase in TAMRA fluorescence at 582 nm. If TAMRA is being excited by energy transfer from quenched 6-FAM, then loss of quenching attributable to hybridization should cause a decrease in the fluorescence emission of TAMRA. The fact that the fluorescence emission of TAMRA increases indicates that the situation is more complex. For example, we have anecdotal evidence that the bases of the oligonucleotide, especially G, quench the fluorescence of both 6-FAM and TAMRA to some degree. When double-stranded, base-pairing may reduce the ability of the bases to quench. The primary factor causing the quenching of 6-FAM in an intact probe is the TAMRA dye. Evidence for the importance of TAMRA is that 6-FAM fluorescence remains relatively unchanged when probes labeled only with 6-FAM are used in the 5' nuclease PCR assay (data not shown). Secondary effectors of fluorescence, both before and after cleavage of the probe, need to be explored further.

Regardless of the physical mechanism, the relative independence of position and quenching greatly simplifies the design of probes for the 5' nuclease PCR assay. There are three main factors that determine the performance of a double-labeled fluorescent probe in the 5' nuclease PCR assay. The first factor is the degree of quenching observed in the intact probe. This is characterized by the value of RQ^- , which is the ratio of reporter to quencher fluorescent emissions for a no template control PCR. Influences on the value of RQ^- include the particular reporter and quencher

TABLE 3 Comparison of Fluorescence Emissions of Single-stranded and Double-stranded Fluorogenic Probes

Probe	518 nm		582 nm		RQ	
	ss	ds	ss	ds	ss	ds
A1-7	27.75	68.53	61.08	138.18	0.45	0.50
A1-26	43.31	509.38	53.50	93.86	0.81	5.43
A3-6	16.75	62.88	39.33	165.57	0.43	0.38
A3-24	30.05	578.64	67.72	140.25	0.45	3.21
P2-7	35.02	70.13	54.63	121.09	0.64	0.58
P2-27	39.89	320.47	65.10	61.13	0.61	5.25
P5-10	27.34	144.85	61.95	165.54	0.44	0.87
P5-28	33.65	462.29	72.39	104.61	0.46	4.43

(ss) Single-stranded. The fluorescence emissions at 518 or 582 nm for solutions containing a final concentration of 50 nM indicated probe, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, and 10 mM $MgCl_2$. (ds) Double-stranded. The solutions contained, in addition, 100 nM A1C for probes A1-7 and A1-26, 100 nM A3C for probes A3-6 and A3-24, 100 nM P2C for probes P2-7 and P2-27, or 100 nM P5C for probes P5-10 and P5-28. Before the addition of $MgCl_2$, 120 μ l of each sample was heated at 95°C for 5 min. Following the addition of 80 μ l of 25 mM $MgCl_2$, each sample was allowed to cool to room temperature and the fluorescence emissions were measured. Reported values are the average of three determinations.

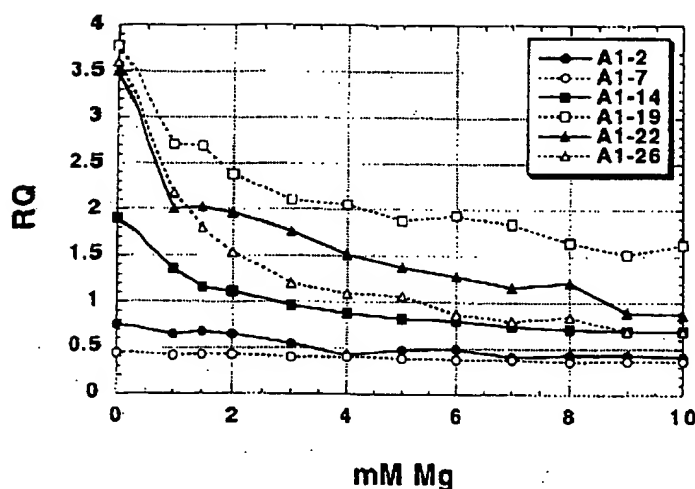


FIGURE 3 Effect of Mg^{2+} concentration on RQ ratio for the A1 series of probes. The fluorescence emission intensity at 518 and 582 nm was measured for solutions containing 50 nM probe, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, and varying amounts (0–10 mM) of $MgCl_2$. The calculated RQ ratios (518 nm intensity divided by 582 nm intensity) are plotted vs. $MgCl_2$ concentration (mM Mg). The key (upper right) shows the probes examined.

dyes used, spacing between reporter and quencher dyes, nucleotide sequence context effects, presence of structure or other factors that reduce flexibility of the oligonucleotide, and purity of the probe. The second factor is the efficiency of hybridization, which depends on probe T_m , presence of secondary structure in probe or template, annealing temperature, and other reaction conditions. The third factor is the efficiency at which *Taq* DNA polymerase cleaves the bound probe between the reporter and quencher dyes. This cleavage is dependent on sequence complementarity between probe and template as shown by the observation that mismatches in the segment between reporter and quencher dyes drastically reduce the cleavage of probe.⁽¹⁾

The rise in RQ^- values for the A1 series of probes seems to indicate that the degree of quenching is reduced somewhat as the quencher is placed toward the 3' end. The lowest apparent quenching is observed for probe A1-19 (see Fig. 3) rather than for the probe where the TAMRA is at the 3' end (A1-26). This is understandable, as the conformation of the 3' end position would be expected to be less restricted than the conformation of an internal position. In effect, a quencher at the 3' end is freer to adopt conformations close to the 5' reporter dye than is an internally placed quencher. For the other three sets of

probes, the interpretation of RQ^- values is less clear-cut. The A3 probes show the same trend as A1, with the 3' TAMRA probe having a larger RQ^- than the internal TAMRA probe. For the P2 pair, both probes have about the same RQ^- value. For the P5 probes, the RQ^- for the 3' probe is less than for the internally labeled probe. Another factor that may explain some of the observed variation is that purity affects the RQ^- value. Although all probes are HPLC purified, a small amount of contamination with unquenched reporter can have a large effect on RQ^- .

Although there may be a modest effect on degree of quenching, the position of the quencher apparently can have a large effect on the efficiency of probe cleavage. The most drastic effect is observed with probe A1-2, where placement of the TAMRA on the second nucleotide reduces the efficiency of cleavage to almost zero. For the A3, P2, and P5 probes, ΔRQ is much greater for the 3' TAMRA probes as compared with the internal TAMRA probes. This is explained most easily by assuming that probes with TAMRA at the 3' end are more likely to be cleaved between reporter and quencher than are probes with TAMRA attached internally. For the A1 probes, the cleavage efficiency of probe A1-7 must already be quite high, as ΔRQ does not increase when the quencher is placed closer to the 3' end. This illus-

trates the importance of being able to use probes with a quencher on the 3' end in the 5' nuclease PCR assay. In this assay, an increase in the intensity of reporter fluorescence is observed only when the probe is cleaved between the reporter and quencher dyes. By placing the reporter and quencher dyes on the opposite ends of an oligonucleotide probe, any cleavage that occurs will be detected. When the quencher is attached to an internal nucleotide, sometimes the probe works well (A1-7) and other times not so well (A3-6). The relatively poor performance of probe A3-6 presumably means the probe is being cleaved 3' to the quencher rather than between the reporter and quencher. Therefore, the best chance of having a probe that reliably detects accumulation of PCR product in the 5' nuclease PCR assay is to use a probe with the reporter and quencher dyes on opposite ends.

Placing the quencher dye on the 3' end may also provide a slight benefit in terms of hybridization efficiency. The presence of a quencher attached to an internal nucleotide might be expected to disrupt base-pairing and reduce the T_m of a probe. In fact, a 2°C–3°C reduction in T_m has been observed for two probes with internally attached TAMRAs.⁽⁹⁾ This disruptive effect would be minimized by placing the quencher at the 3' end. Thus, probes with 3' quenchers might exhibit slightly higher hybridization efficiencies than probes with internal quenchers.

The combination of increased cleavage and hybridization efficiencies means that probes with 3' quenchers probably will be more tolerant of mismatches between probe and target as compared with internally labeled probes. This tolerance of mismatches can be advantageous, as when trying to use a single probe to detect PCR-amplified products from samples of different species. Also, it means that cleavage of probe during PCR is less sensitive to alterations in annealing temperature or other reaction conditions. The one application where tolerance of mismatches may be a disadvantage is for allelic discrimination. Lee et al.⁽¹¹⁾ demonstrated that allele-specific probes were cleaved between reporter and quencher only when hybridized to a perfectly complementary target. This allowed them to distinguish the normal human cystic fibrosis allele from the $\Delta F508$ mutant. Their probes had TAMRA attached to the seventh nucleotide from

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the 5' end and were designed so that any mismatches were between the reporter and quencher. Increasing the distance between reporter and quencher would lessen the disruptive effect of mismatches and allow cleavage of the probe on the incorrect target. Thus, probes with a quencher attached to an internal nucleotide may still be useful for allelic discrimination.

In this study loss of quenching upon hybridization was used to show that quenching by a 3' TAMRA is dependent on the flexibility of a single-stranded oligonucleotide. The increase in reporter fluorescence intensity, though, could also be used to determine whether hybridization has occurred or not. Thus, oligonucleotides with reporter and quencher dyes attached at opposite ends should also be useful as hybridization probes. The ability to detect hybridization in real time means that these probes could be used to measure hybridization kinetics. Also, this type of probe could be used to develop homogeneous hybridization assays for diagnostics or other applications. Bagwell et al.⁽¹⁰⁾ describe just this type of homogeneous assay where hybridization of a probe causes an increase in fluorescence caused by a loss of quenching. However, they utilized a complex probe design that requires adding nucleotides to both ends of the probe sequence to form two imperfect hairpins. The results presented here demonstrate that the simple addition of a reporter dye to one end of an oligonucleotide and a quencher dye to the other end generates a fluorogenic probe that can detect hybridization or PCR amplification.

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GENOME METHODS

Real Time Quantitative PCR

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We have developed a novel "real time" quantitative PCR method. The method measures PCR product accumulation through a dual-labeled fluorogenic probe (i.e., TaqMan Probe). This method provides very accurate and reproducible quantitation of gene copies. Unlike other quantitative PCR methods, real-time PCR does not require post-PCR sample handling, preventing potential PCR product carry-over contamination and resulting in much faster and higher throughput assays. The real-time PCR method has a very large dynamic range of starting target molecule determination (at least five orders of magnitude). Real-time quantitative PCR is extremely accurate and less labor-intensive than current quantitative PCR methods.

Quantitative nucleic acid sequence analysis has had an important role in many fields of biological research. Measurement of gene expression (RNA) has been used extensively in monitoring biological responses to various stimuli (Tan et al. 1994; Huang et al. 1995a,b; Prud'homme et al. 1995). Quantitative gene analysis (DNA) has been used to determine the genome quantity of a particular gene, as in the case of the human *HER2* gene, which is amplified in ~30% of breast tumors (Slamon et al. 1987). Gene and genome quantitation (DNA and RNA) also have been used for analysis of human immunodeficiency virus (HIV) burden demonstrating changes in the levels of virus throughout the different phases of the disease (Connor et al. 1993; Platak et al. 1993b; Furtado et al. 1995).

Many methods have been described for the quantitative analysis of nucleic acid sequences (both for RNA and DNA; Southern 1975; Sharp et al. 1980; Thomas 1980). Recently, PCR has proven to be a powerful tool for quantitative nucleic acid analysis. PCR and reverse transcriptase (RT)-PCR have permitted the analysis of minimal starting quantities of nucleic acid (as little as one cell equivalent). This has made possible many experiments that could not have been performed with traditional methods. Although PCR has provided a powerful tool, it is imperative

that it be used properly for quantitation (Rauymaekers 1995). Many early reports of quantitative PCR and RT-PCR described quantitation of the PCR product but did not measure the initial target sequence quantity. It is essential to design proper controls for the quantitation of the initial target sequences (Herre 1992; Clementi et al. 1993).

Researchers have developed several methods of quantitative PCR and RT-PCR. One approach measures PCR product quantity in the log phase of the reaction before the plateau (Kellogg et al. 1990; Pang et al. 1990). This method requires that each sample has equal input amounts of nucleic acid and that each sample under analysis amplifies with identical efficiency up to the point of quantitative analysis. A gene sequence (contained in all samples at relatively constant quantities, such as β -actin) can be used for sample amplification efficiency normalization. Using conventional methods of PCR detection and quantitation (gel electrophoresis or plate capture hybridization), it is extremely laborious to assure that all samples are analyzed during the log phase of the reaction (for both the target gene and the normalization gene). Another method, quantitative competitive (QC)-PCR, has been developed and is used widely for PCR quantitation. QC-PCR relies on the inclusion of an internal control competitor in each reaction (Becker-Andre 1991; Platak et al. 1993a,b). The efficiency of each reaction is normalized to the internal competitor. A known amount of internal competitor can be

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added to each sample. To obtain relative quantitation, the unknown target PCR product is compared with the known competitor PCR product. Success of a quantitative competitive PCR assay relies on developing an internal control that amplifies with the same efficiency as the target molecule. The design of the competitor and the validation of amplification efficiencies require a dedicated effort. However, because QC-PCR does not require that PCR products be analyzed during the log phase of the amplification, it is the easier of the two methods to use.

Several detection systems are used for quantitative PCR and RT-PCR analysis: (1) agarose gels, (2) fluorescent labeling of PCR products and detection with laser-induced fluorescence using capillary electrophoresis (Fusco et al. 1995; Williams et al. 1996) or acrylamide gels, and (3) plate capture and sandwich probe hybridization (Mulder et al. 1994). Although these methods proved successful, each method requires post-PCR manipulations that add time to the analysis and may lead to laboratory contamination. The sample throughput of these methods is limited (with the exception of the plate capture approach), and, therefore, these methods are not well suited for uses demanding high sample throughput (i.e., screening of large numbers of biomolecules or analyzing samples for diagnostics or clinical trials).

Here we report the development of a novel assay for quantitative DNA analysis. The assay is based on the use of the 5' nuclease assay first described by Holland et al. (1991). The method uses the 5' nuclease activity of *Taq* polymerase to cleave a nonextendible hybridization probe during the extension phase of PCR. The approach uses dual-labeled fluorogenic hybridization probes (Lee et al. 1993; Bassler et al. 1995; Livak et al. 1995a,b). One fluorescent dye serves as a reporter (FAM (i.e., 6-carboxyfluorescein)) and its emission spectra is quenched by the second fluorescent dye, TAMRA (i.e., 6-carboxy-tetramethylrhodamine). The nuclease degradation of the hybridization probe releases the quenching of the FAM fluorescent emission, resulting in an increase in peak fluorescent emission at 518 nm. The use of a sequence detector (ABI Prism) allows measurement of fluorescent spectra of all 96 wells of the thermal cycler continuously during the PCR amplification. Therefore, the reactions are monitored in real time. The output data is described and quantitative analysis of input target DNA sequences is discussed below.

RESULTS

PCR Product Detection in Real Time

The goal was to develop a high-throughput, sensitive, and accurate gene quantitation assay for use in monitoring lipid mediated therapeutic gene delivery. A plasmid encoding human factor VIII gene sequence, pF8TM (see Methods), was used as a model therapeutic gene. The assay uses fluorescent Taqman methodology and an instrument capable of measuring fluorescence in real time (ABI Prism 7700 Sequence Detector). The Taqman reaction requires a hybridization probe labeled with two different fluorescent dyes. One dye is a reporter dye (FAM), the other is a quenching dye (TAMRA). When the probe is intact, fluorescent energy transfer occurs and the reporter dye fluorescent emission is absorbed by the quenching dye (TAMRA). During the extension phase of the PCR cycle, the fluorescent hybridization probe is cleaved by the 5'-3' nucleolytic activity of the DNA polymerase. On cleavage of the probe, the reporter dye emission is no longer transferred efficiently to the quenching dye, resulting in an increase of the reporter dye fluorescent emission spectra. PCR primers and probes were designed for the human factor VIII sequence and human β -actin gene (as described in Methods). Optimization reactions were performed to choose the appropriate probe and magnesium concentrations yielding the highest intensity of reporter fluorescent signal without sacrificing specificity. The instrument uses a charge-coupled device (i.e., CCD camera) for measuring the fluorescent emission spectra from 500 to 650 nm. Each PCR tube was monitored sequentially for 25 msec with continuous monitoring throughout the amplification. Each tube was re-examined every 8.5 sec. Computer software was designed to examine the fluorescent intensity of both the reporter dye (FAM) and the quenching dye (TAMRA). The fluorescent intensity of the quenching dye, TAMRA, changes very little over the course of the PCR amplification (data not shown). Therefore, the intensity of TAMRA dye emission serves as an internal standard with which to normalize the reporter dye (FAM) emission variations. The software calculates a value termed ΔRn (or $\Delta R(2)$) using the following equation: $\Delta Rn = (Rn^+ - Rn^-) / Rn^-$, where Rn^+ = emission intensity of reporter/emission intensity of quencher at any given time in a reaction tube, and Rn^- = emission intensity of re-

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porter/emission intensity of quencher measured prior to PCR amplification in that same reaction tube. For the purpose of quantitation, the last three data points (ΔRn s) collected during the extension step for each PCR cycle were analyzed. The nucleolytic degradation of the hybridization probe occurs during the extension phase of PCR, and, therefore, reporter fluorescent emission increases during this time. The three data points were averaged for each PCR cycle and the mean value for each was plotted in an "amplification plot" shown in Figure 1A. The ΔRn mean value is plotted on the y-axis, and time, represented by cycle number, is plotted on the x-axis. During the early cycles of the PCR amplification, the ΔRn

value remains at base line. When sufficient hybridization probe has been cleaved by the *Taq* polymerase nuclease activity, the intensity of reporter fluorescent emission increases. Most PCR amplifications reach a plateau phase of reporter fluorescent emission if the reaction is carried out to high cycle numbers. The amplification plot is examined early in the reaction, at a point that represents the log phase of product accumulation. This is done by assigning an arbitrary threshold that is based on the variability of the base-line data. In Figure 1A, the threshold was set at 10 standard deviations above the mean of base line emission calculated from cycles 1 to 15. Once the threshold is chosen, the point at which

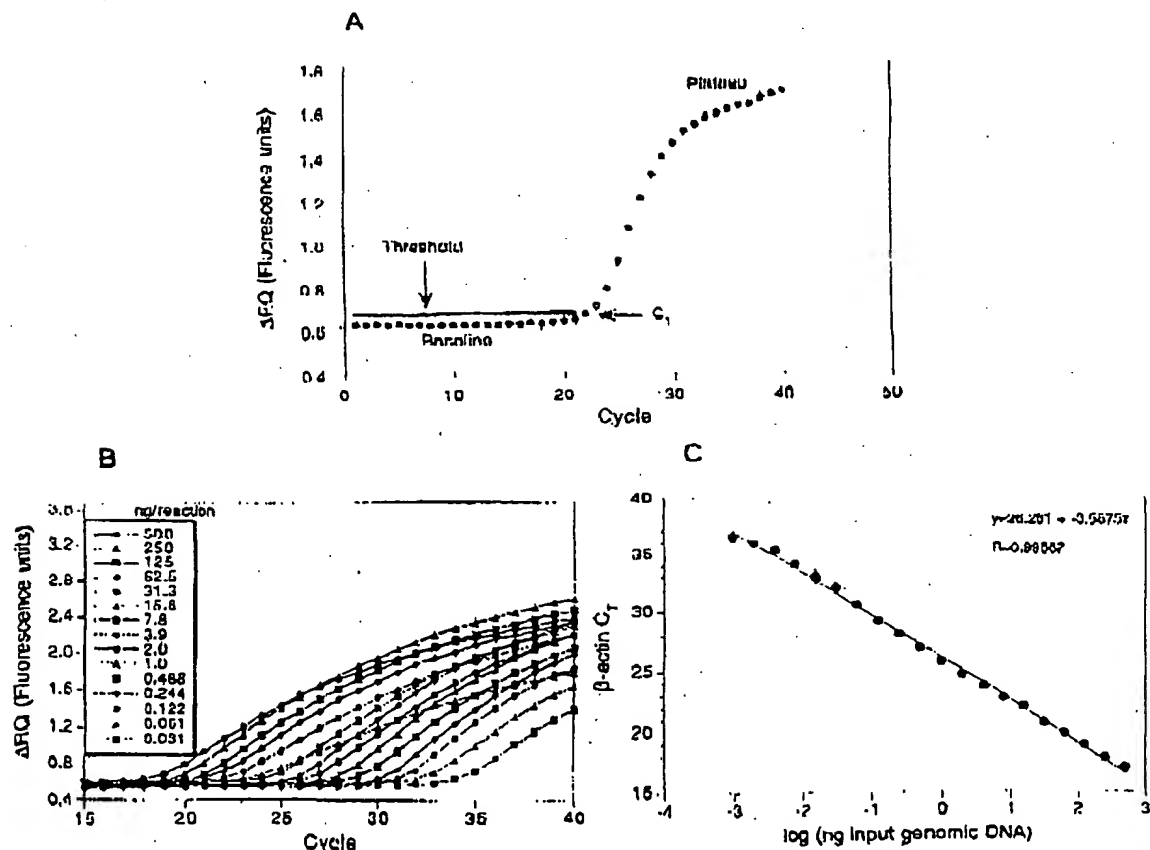


Figure 1 PCR product detection in real time. (A) The Model 7700 software will construct amplification plots from the extension phase fluorescent emission data collected during the PCR amplification. The standard deviation is determined from the data points collected from the base line of the amplification plot. C_T values are calculated by determining the point at which the fluorescence exceeds a threshold limit (usually 10 times the standard deviation of the base line). (B) Overlay of amplification plots of serially (1:2) diluted human genomic DNA samples amplified with β -actin primers. (C) Input DNA concentration of the samples plotted versus C_T . All

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the amplification plot crosses the threshold is defined as C_T . C_T is reported as the cycle number at this point. As will be demonstrated, the C_T value is predictive of the quantity of input target.

C_T Values Provide a Quantitative Measurement of Input Target Sequences

Figure 1B shows amplification plots of 15 different PCR amplifications overlaid. The amplifications were performed on a 1:2 serial dilution of human genomic DNA. The amplified target was human β actin. The amplification plots shift to the right (to higher threshold cycles) as the input target quantity is reduced. This is expected because reactions with fewer starting copies of the target molecule require greater amplification to degrade enough probe to attain the threshold fluorescence. An arbitrary threshold of 10 standard deviations above the base line was used to determine the C_T values. Figure 1C represents the C_T values plotted versus the sample dilution value. Each dilution was amplified in triplicate PCR amplifications and plotted as mean values with error bars representing one standard deviation. The C_T values decrease linearly with increasing target quantity. Thus, C_T values can be used as a quantitative measurement of the input target number. It should be noted that the amplification plot for the 15.6-ng sample shown in Figure 1B does not reflect the same fluorescent rate of increase exhibited by most of the other samples. The 15.6-ng sample also achieves endpoint plateau at a lower fluorescent value than would be expected based on the input DNA. This phenomenon has been observed occasionally with other samples (data not shown) and may be attributable to late cycle inhibition; this hypothesis is still under investigation. It is important to note that the flattened slope and early plateau do not impact significantly the calculated C_T value as demonstrated by the fit on the line shown in Figure 1C. All triplicate amplifications resulted in very similar C_T values—the standard deviation did not exceed 0.5 for any dilution. This experiment contains a >100,000-fold range of input target molecules. Using C_T values for quantitation permits a much larger assay range than directly using total fluorescent emission intensity for quantitation. The linear range of fluorescent intensity measurement of the ABI Prism 7700 Se-

ments over a very large range of relative starting target quantities.

Sample Preparation Validation

Several parameters influence the efficiency of PCR amplification: magnesium and salt concentrations, reaction conditions (i.e., time and temperature), PCR target size and composition, primer sequences, and sample purity. All of the above factors are common to a single PCR assay except sample to sample purity. In an effort to validate the method of sample preparation for the factor VIII assay, PCR amplification reproducibility and efficiency of 10 replicate sample preparations were examined. After genomic DNA was prepared from the 10 replicate samples, the DNA was quantitated by ultraviolet spectroscopy. Amplifications were performed analyzing β -actin gene content in 100 and 25 ng of total genomic DNA. Each PCR amplification was performed in triplicate. Comparison of C_T values for each triplicate sample show minimal variation based on standard deviation and coefficient of variance (Table 1). Therefore, each of the triplicate PCR amplifications was highly reproducible, demonstrating that real time PCR using this instrumentation introduces minimal variation into the quantitative PCR analysis. Comparison of the mean C_T values of the 10 replicate sample preparations also showed minimal variability, indicating that each sample preparation yielded similar results for β -actin gene quantity. The highest C_T difference between any of the samples was 0.8 and 0.71 for the 100 and 25 ng samples, respectively. Additionally, the amplification of each sample exhibited an equivalent rate of fluorescent emission intensity change per amount of DNA target analyzed as indicated by similar slopes derived from the sample dilutions (Fig. 2). Any sample containing an excess of a PCR inhibitor would exhibit a greater measured β -actin C_T value for a given quantity of DNA. In addition, the inhibitor would be diluted along with the sample in the dilution analysis (Fig. 2), altering the expected C_T value change. Each sample amplification yielded a similar result in the analysis, demonstrating that this method of sample preparation is highly reproducible with regard to sample purity.

Quantitative Analysis of a Plasmid After

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Table 1. Reproducibility of Sample Preparation Method

Sample no.	100 ng				25 ng			
	C _T	mean	standard deviation	CV	C _T	mean	standard deviation	CV
1	18.24	18.27	0.06	0.32	20.48	20.51	0.03	0.17
	18.23				20.55			
	18.33				20.5			
2	18.33	18.37	0.06	0.32	20.61	20.54	0.11	0.54
	18.35				20.59			
	18.44				20.41			
3	18.3	18.34	0.07	0.36	20.54	20.54	0.06	0.28
	18.3				20.6			
	18.42				20.49			
4	18.15	18.23	0.08	0.46	20.48	20.43	0.05	0.26
	18.23				20.44			
	18.32				20.38			
5	18.4	18.42	0.04	0.23	20.68	20.73	0.13	0.61
	18.38				20.87			
	18.46				20.63			
6	18.54	18.71	0.24	1.26	21.09	21.06	0.03	0.15
	18.67				21.04			
	19				21.01			
7	18.28	18.39	0.12	0.66	20.67	20.68	0.04	0.2
	18.36				20.73			
	18.52				20.65			
8	18.45	18.63	0.16	0.83	20.98	20.86	0.12	0.57
	18.7				20.84			
	18.73				20.75			
9	18.18	18.29	0.1	0.55	20.46	20.51	0.07	0.32
	18.34				20.54			
	18.26				20.48			
10	18.42	18.55	0.12	0.65	20.79	20.73	0.1	0.16
	18.57				20.78			
	18.66				20.62			
Mean	(1 10)	18.42	0.17	0.90		20.66	0.19	0.94

for containing a partial cDNA for human factor VIII, pF8TM. A series of transfections was set up using a decreasing amount of the plasmid (40, 4, 0.5, and 0.1 µg). Twenty-four hours post-transfection, total DNA was purified from each flask of cells. β -Actin gene quantity was chosen as a value for normalization of genomic DNA concentration from each sample. In this experiment, β -actin gene content should remain constant relative to total genomic DNA. Figure 3 shows the result of the β -actin DNA measurement (100 ng total DNA determined by ultraviolet spectroscopy) of each sample. Each sample was analyzed in triplicate and the mean β -actin C_T values of the triplicates were plotted (error bars represent one standard deviation). The highest difference

between any two sample means was 0.95 C_T. Ten nanograms of total DNA of each sample were also examined for β -actin. The results again showed that very similar amounts of genomic DNA were present; the maximum mean β -actin C_T value difference was 1.0. As Figure 3 shows, the rate of β -actin C_T change between the 100 and 10-ng samples was similar (slope values range between 3.56 and -3.45). This verifies again that the method of sample preparation yields samples of identical PCR integrity (i.e., no sample contained an excessive amount of a PCR inhibitor). However, these results indicate that each sample contained slight differences in the actual amount of genomic DNA analyzed. Determination of actual genomic DNA concentration was accomplished

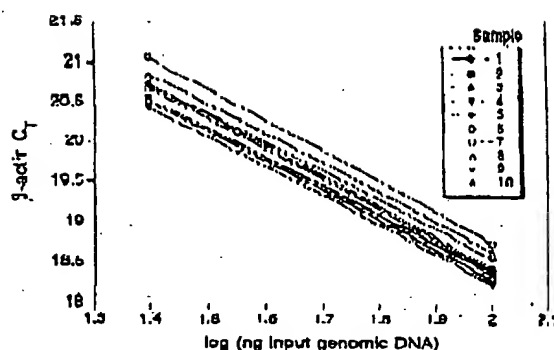


Figure 2 Sample preparation purity. The replicate samples shown in Table 1 were also amplified in triplicate using 25 ng of each DNA sample. The figure shows the input DNA concentration (100 and 25 ng) vs. C_t . In the figure, the 100 and 25 ng points for each sample are connected by a line.

by plotting the mean β -actin C_t value obtained for each 100-ng sample on a β -actin standard curve (shown in Fig. 4C). The actual genomic DNA concentration of each sample, a , was obtained by extrapolation to the x-axis.

Figure 4A shows the measured (i.e., non-normalized) quantities of factor VIII plasmid DNA (pF8TM) from each of the four transient cell transfections. Each reaction contained 100 ng of total sample DNA (as determined by UV spectroscopy). Each sample was analyzed in triplicate

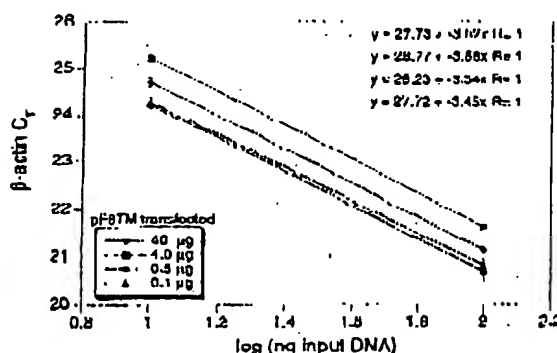


Figure 3 Analysis of transfected cell DNA quantity and purity. The DNA preparations of the four 293 cell transfections (40, 4, 0.5, and 0.1 μ g of pF8TM) were analyzed for the β -actin gene. 100 and 10 ng (determined by ultraviolet spectroscopy) of each sample were amplified in triplicate. For each amount of pF8TM that was transfected, the β -actin C_t values are plotted versus the total input DNA concentration.

REAL TIME-QUANTITATIVE PCR

PCR amplifications. As shown, pF8TM purified from the 293 cells decreases (mean C_t values increase) with decreasing amounts of plasmid (transfected). The mean C_t values obtained for pF8TM in Figure 4A were plotted on a standard curve comprised of serially diluted pF8TM shown in Figure 4B. The quantity of pF8TM, b found in each of the four transfections was determined by extrapolation to the x-axis of the standard curve in Figure 4B. These uncorrected values, b , for pF8TM were normalized to determine the actual amount of pF8TM found per 100 ng of genomic DNA by using the equation:

$$\frac{b \times 100 \text{ ng}}{a} = \text{actual pF8TM copies per 100 ng of genomic DNA}$$

where a = actual genomic DNA in a sample and b = pF8TM copies from the standard curve. The normalized quantity of pF8TM per 100 ng of genomic DNA for each of the four transfections is shown in Figure 4D. These results show that the quantity of factor VIII plasmid associated with the 293 cells, 24 hr after transfection, decreases with decreasing plasmid concentration used in the transfection. The quantity of pF8TM associated with 293 cells, after transfection with 40 μ g of plasmid, was 35 pg per 100 ng genomic DNA. This results in ~520 plasmid copies per cell.

DISCUSSION

We have described a new method for quantitating gene copy numbers using real-time analysis of PCR amplifications. Real-time PCR is compatible with either of the two PCR (RT-PCR) approaches: (1) quantitative competitive where an internal competitor for each target sequence is used for normalization (data not shown) or (2) quantitative comparative PCR using a normalization gene contained within the sample (i.e., β -actin) or a "housekeeping" gene for RT-PCR. If equal amounts of nucleic acid are analyzed for each sample and if the amplification efficiency before quantitative analysis is identical for each sample, the internal control (normalization gene or competitor) should give equal signals for all samples.

The real-time PCR method offers several advantages over the other two methods currently employed (see the Introduction). First, the real-time PCR method is performed in a closed-tube system and requires no post-PCR manipulation

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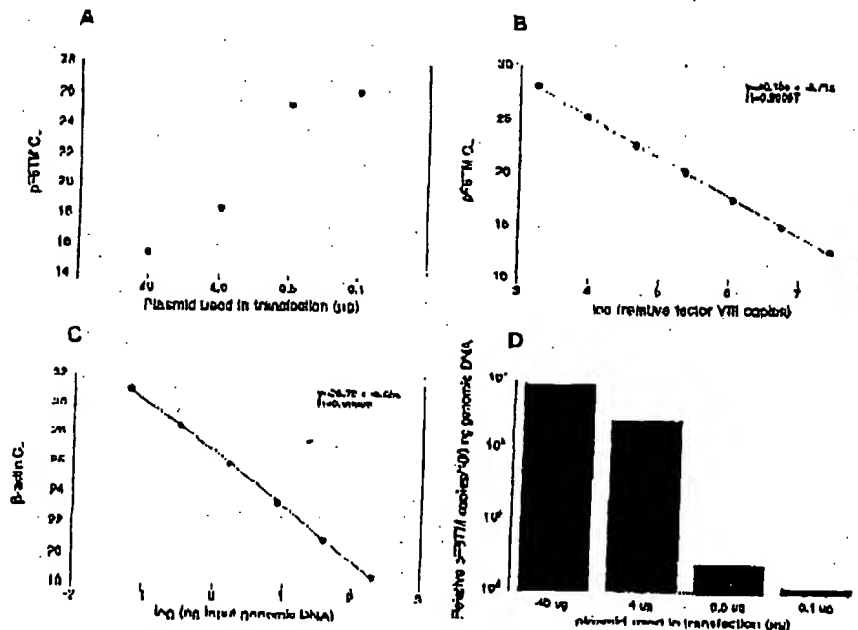


Figure 4 Quantitative analysis of pF8TM in transfected cells. (A) Amount of plasmid DNA used for the transfection plotted against the mean C_T value determined for pF8TM remaining 24 hr after transfection. (B, C) Standard curves of pF8TM and β-actin, respectively. pF8TM DNA (B) and genomic DNA (C) were diluted serially 1:5 before amplification with the appropriate primers. The β-actin standard curve was used to normalize the results of 1 to 100 ng of genomic DNA. (D) The amount of pF8TM present per 100 ng of genomic DNA.

of sample. Therefore, the potential for PCR contamination in the laboratory is reduced because amplified products can be analyzed and disposed of without opening the reaction tubes. Second, this method supports the use of a normalization gene (i.e., β-actin) for quantitative PCR or house-keeping genes for quantitative RT-PCR controls. Analysis is performed in real time during the log phase of product accumulation. Analysis during log phase permits many different genes (over a wide input target range) to be analyzed simultaneously, without concern of reaching reaction plateau at different cycles. This will make multi-gene analysis assays much easier to develop, because individual internal competitors will not be needed for each gene under analysis. Third, sample throughput will increase dramatically with the new method because there is no post-PCR processing time. Additionally, working in a 96-well format is highly compatible with automation technology.

The real-time PCR method is highly reproducible. Replicate amplifications can be analyzed

for each sample minimizing potential error. The system allows for a very large assay dynamic range (approaching 1,000,000-fold starting target). Using a standard curve for the target of interest, relative copy number values can be determined for any unknown sample. Fluorescent threshold values, C_{TP} correlate linearly with relative DNA copy numbers. Real time quantitative RT-PCR methodology (Gibson et al., this issue) has also been developed. Finally, real time quantitative PCR methodology can be used to develop high-throughput screening assays for a variety of applications [quantitative gene expression (RT-PCR), gene copy assays (Her2, HIV, etc.), genotyping (knockout mouse analysis), and immun-

PCR]. Real-time PCR may also be performed using intercalating dyes (Higuchi et al. 1992) such as ethidium bromide. The fluorogenic probe method offers a major advantage over intercalating dyes--greater specificity (i.e., primer dimers and nonspecific PCR products are not detected).

REAL TIME QUANTITATIVE PCR

METHODS

Generation of a Plasmid Containing a Partial cDNA for Human Factor VIII

Total RNA was harvested (RNeasy B from Tel Test, Inc., Friendswood, TX) from cells transfected with a factor VIII expression vector, pCIS2.8c251 (Eaton et al. 1986; Gorman et al. 1990). A factor VIII partial cDNA sequence was generated by RT-PCR (GeneAmp 12.7th RNA PCR Kit (part N808-0179, PE Applied Biosystems, Foster City, CA)) using the PCR primers F8for and F8rev (primer sequences are shown below). The amplicon was reamplified using modified F8for and F8rev primers (appended with *Hind*III and *Hind*III restriction site sequences at the 5' ends) and cloned into pCIS2.32 (Promega Corp., Madison, WI). The resulting clone, pF8TM, was used for transient transfection of 293 cells.

Amplification of Target DNA and Detection of Amplicon Factor VIII Plasmid DNA

(pF8TM) was amplified with the primers F8for 5'-CCGCTTCCCAAGGAGTGAATGTC-3' and F8rev 5'-AAACCTTCCACCTGGATGCTACG-3'. The reaction produced a 422-bp PCR product. The forward primer was designed to recognize a unique sequence found in the 5' untranslated region of the parent pCIS2.8c251 plasmid and therefore does not recognize and amplify the human factor VIII gene. Primers were chosen with the assistance of the computer program Oligo 4.0 (National Biosciences, Inc., Plymouth, MN). The human β -actin gene was amplified with the primers β -actin forward primer 5'-TCACCCACACTCTTCCCATCTTACCA-3' and β -actin reverse primer 5'-CAACCGGAACCCGCTCATTCCTCAATGG-3'. The reaction produced a 295-bp PCR product.

Amplification reactions (50 μ l) contained a DNA sample, 10 \times PCR Buffer II (5 μ l), 200 μ M dATP, dCTP, dGTP, and 400 μ M dUTP, 4 mM MgCl₂, 1.25 Units AmpliTaq DNA polymerase, 0.5 unit AmpliTaq uracil N-glycosylase (UNG), 60 pmole of each factor VIII primer, and 15 pmole of each β -actin primer. The reactions also contained one of the following detection probes (100 nM each): F8probe 5'-(FAM)ACCTCTTCCACCTGCTTCTTCTCTTGCCTT(TAMRA)p-3' and β -actin probe 5'-(FAM)ATGCCXCCX(TAMRA)CCCCCATGCCATCp-3' where p indicates phosphorylation and X indicates a linker arm nucleotide. Reaction tubes were MicroAmp Optical Tubes (part number N801 0933, Perkin Elmer) that were frosted (at Perkin Elmer) to prevent light from reflecting. Tube caps were similar to MicroAmp Caps but specially designed to prevent light scattering. All of the PCR consumables were supplied by PE Applied Biosystems (Foster City, CA) except the factor VIII primers, which were synthesized at Genentech, Inc. (South San Francisco, CA). Probes were designed using the Oligo 4.0 software, following guidelines suggested in the Model 7700 Sequence Detector instrument manual. Briefly, probe T_m should be at least 5 $^{\circ}$ C higher than the annealing temperature used during thermal cycling; primers should not form stable duplexes with the probe.

The thermal cycling conditions included 2 min at 50 $^{\circ}$ C and 10 min at 95 $^{\circ}$ C. Thermal cycling proceeded with

reactions were performed in the Model 7700 Sequence Detector (PE Applied Biosystems), which contains a GeneAmp PCR System 9600. Reaction conditions were programmed on a Power Macintosh 7100 (Apple Computer, Santa Clara, CA) linked directly to the Model 7700 Sequence Detector. Analysis of data was also performed on the Macintosh computer. Collection and analysis software was developed at PE Applied Biosystems.

Transfection of Cells with Factor VIII Construct

Four T175 flasks of 293 cells (ATCC CRL 1573), a human fetal kidney suspension cell line, were grown to 80% confluency and transfected pF8TM. Cells were grown in the following media: 50% HAM'S F12 without GHT, 50% low glucose Dulbecco's modified Eagle medium (DMEM) without glycine with sodium bicarbonate, 10% fetal bovine serum, 2 mM L-glutamine, and 1% penicillin-streptomycin. The media was changed 30 min before the transfection. pF8TM DNA amounts of 40, 4, 0.5, and 0.1 μ g were added to 1.5 ml of a solution containing 0.125 M CaCl₂ and 1 \times HEPES. The four mixtures were left at room temperature for 10 min and then added dropwise to the cells. The flasks were incubated at 37 $^{\circ}$ C and 5% CO₂ for 24 hr washed with PBS, and resuspended in PBS. The resuspended cells were divided into aliquots and DNA was extracted immediately using the QIAamp Blood Kit (Qiagen Crawsforth, CA). DNA was eluted into 200 μ l of 20 mM Tris-HCl at pH 8.0.

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WISP genes are members of the connective tissue growth factor family that are up-regulated in Wnt-1-transformed cells and aberrantly expressed in human colon tumors

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ABSTRACT Wnt family members are critical to many developmental processes, and components of the Wnt signaling pathway have been linked to tumorigenesis in familial and sporadic colon carcinomas. Here we report the identification of two genes, *WISP-1* and *WISP-2*, that are up-regulated in the mouse mammary epithelial cell line C57MG transformed by Wnt-1, but not by Wnt-4. Together with a third related gene, *WISP-3*, these proteins define a subfamily of the connective tissue growth factor family. Two distinct systems demonstrated *WISP* induction to be associated with the expression of Wnt-1. These included (i) C57MG cells infected with a Wnt-1 retroviral vector or expressing Wnt-1 under the control of a tetracycline repressible promoter, and (ii) Wnt-1 transgenic mice. The *WISP-1* gene was localized to human chromosome 8q24.1-8q24.3. *WISP-1* genomic DNA was amplified in colon cancer cell lines and in human colon tumors and its RNA overexpressed (2- to >30-fold) in 84% of the tumors examined compared with patient-matched normal mucosa. *WISP-3* mapped to chromosome 6q22-6q23 and also was overexpressed (4- to >40-fold) in 63% of the colon tumors analyzed. In contrast, *WISP-2* mapped to human chromosome 20q12-20q13 and its DNA was amplified, but RNA expression was reduced (2- to >30-fold) in 79% of the tumors. These results suggest that the *WISP* genes may be downstream of Wnt-1 signaling and that aberrant levels of *WISP* expression in colon cancer may play a role in colon tumorigenesis.

Wnt-1 is a member of an expanding family of cysteine-rich, glycosylated signaling proteins that mediate diverse developmental processes such as the control of cell proliferation, adhesion, cell polarity, and the establishment of cell fates (1, 2). Wnt-1 originally was identified as an oncogene activated by the insertion of mouse mammary tumor virus in virus-induced mammary adenocarcinomas (3, 4). Although Wnt-1 is not expressed in the normal mammary gland, expression of Wnt-1 in transgenic mice causes mammary tumors (5).

In mammalian cells, Wnt family members initiate signaling by binding to the seven-transmembrane spanning Frizzled receptors and recruiting the cytoplasmic protein Dishevelled (Dsh) to the cell membrane (1, 2, 6). Dsh then inhibits the kinase activity of the normally constitutively active glycogen synthase kinase-3 β (GSK-3 β) resulting in an increase in β -catenin levels. Stabilized β -catenin interacts with the transcription factor TCF/Lef1, forming a complex that appears in

the nucleus and binds TCF/Lef1 target DNA elements to activate transcription (7, 8). Other experiments suggest that the adenomatous polyposis coli (APC) tumor suppressor gene also plays an important role in Wnt signaling by regulating β -catenin levels (9). APC is phosphorylated by GSK-3 β , binds to β -catenin, and facilitates its degradation. Mutations in either APC or β -catenin have been associated with colon carcinomas and melanomas, suggesting these mutations contribute to the development of these types of cancer, implicating the Wnt pathway in tumorigenesis (1).

Although much has been learned about the Wnt signaling pathway over the past several years, only a few of the transcriptionally activated downstream components activated by Wnt have been characterized. Those that have been described cannot account for all of the diverse functions attributed to Wnt signaling. Among the candidate Wnt target genes are those encoding the nodal-related 3 gene, *Xnr3*, a member of the transforming growth factor (TGF)- β superfamily, and the homeobox genes, *engrailed*, *gooseoid*, *twin* (*Xtwn*), and *siamois* (2). A recent report also identifies *c-myc* as a target gene of the Wnt signaling pathway (10).

To identify additional downstream genes in the Wnt signaling pathway that are relevant to the transformed cell phenotype, we used a PCR-based cDNA subtraction strategy, suppression subtractive hybridization (SSH) (11), using RNA isolated from C57MG mouse mammary epithelial cells and C57MG cells stably transformed by a Wnt-1 retrovirus. Overexpression of Wnt-1 in this cell line is sufficient to induce a partially transformed phenotype, characterized by elongated and refractile cells that lose contact inhibition and form a multilayered array (12, 13). We reasoned that genes differentially expressed between these two cell lines might contribute to the transformed phenotype.

In this paper, we describe the cloning and characterization of two genes up-regulated in Wnt-1 transformed cells, *WISP-1* and *WISP-2*, and a third related gene, *WISP-3*. The *WISP* genes are members of the CCN family of growth factors, which includes connective tissue growth factor (CTGF), Cyr61, and nov, a family not previously linked to Wnt signaling.

MATERIALS AND METHODS

SSH. SSH was performed by using the PCR-Select cDNA Subtraction Kit (CLONTECH). Tester double-stranded

Abbreviations: TGF, transforming growth factor; CTGF, connective tissue growth factor; SSH, suppression subtractive hybridization; VWC, von Willebrand factor type C module.

Data deposition: The sequences reported in this paper have been deposited in the Genbank database (accession nos. AF100777, AF100778, AF100779, AF100780, and AF100781).

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cDNA was synthesized from 2 μ g of poly(A)⁺ RNA isolated from the C57MG/Wnt-1 cell line and driver cDNA from 2 μ g of poly(A)⁺ RNA from the parent C57MG cells. The subtracted cDNA library was subcloned into a pGEM-T vector for further analysis.

cDNA Library Screening. Clones encoding full-length mouse *WISP-1* were isolated by screening a λ gt10 mouse embryo cDNA library (CLONTECH) with a 70-bp probe from the original partial clone 568 sequence corresponding to amino acids 128–169. Clones encoding full-length human *WISP-1* were isolated by screening λ gt10 lung and fetal kidney cDNA libraries with the same probe at low stringency. Clones encoding full-length mouse and human *WISP-2* were isolated by screening a C57MG/Wnt-1 or human fetal lung cDNA library with a probe corresponding to nucleotides 1463–1512. Full-length cDNAs encoding *WISP-3* were cloned from human bone marrow and fetal kidney libraries.

Expression of Human *WISP* RNA. PCR amplification of first-strand cDNA was performed with human Multiple Tissue cDNA panels (CLONTECH) and 300 μ M of each dNTP at 94°C for 1 sec, 62°C for 30 sec, 72°C for 1 min, for 22–32 cycles. *WISP* and glyceraldehyde-3-phosphate dehydrogenase primer sequences are available on request.

In Situ Hybridization. ³²P-labeled sense and antisense riboprobes were transcribed from an 897-bp PCR product corresponding to nucleotides 601–1440 of mouse *WISP-1* or a 294-bp PCR product corresponding to nucleotides 82–375 of mouse *WISP-2*. All tissues were processed as described (40).

Radiation Hybrid Mapping. Genomic DNA from each hybrid in the Stanford G3 and Genebridge4 Radiation Hybrid Panels (Research Genetics, Huntsville, AL) and human and hamster control DNAs were PCR-amplified, and the results were submitted to the Stanford or Massachusetts Institute of Technology web servers.

Cell Lines, Tumors, and Mucosa Specimens. Tissue specimens were obtained from the Department of Pathology (University of Pittsburgh) for patients undergoing colon resection and from the University of Leeds, United Kingdom. Genomic DNA was isolated (Qiagen) from the pooled blood of 10 normal human donors, surgical specimens, and the following ATCC human cell lines: SW480, COLO 320DM, HT-29, WiDr, and SW403 (colon adenocarcinomas), SW620 (lymph node metastasis, colon adenocarcinoma), HCT 116 (colon carcinoma), SK-CO-1 (colon adenocarcinoma, ascites), and HM7 (a variant of ATCC colon adenocarcinoma cell line LS 174T). DNA concentration was determined by using Hoechst dye 33258 intercalation fluorimetry. Total RNA was prepared by homogenization in 7 M GuSCN followed by centrifugation over CsCl cushions or prepared by using RNeasy.

Gene Amplification and RNA Expression Analysis. Relative gene amplification and RNA expression of *WISPs* and *c-myc* in the cell lines, colorectal tumors, and normal mucosa were determined by quantitative PCR. Gene-specific primers and fluorogenic probes (sequences available on request) were designed and used to amplify and quantitate the genes. The relative gene copy number was derived by using the formula $2^{\Delta\Delta C_t}$ where ΔC_t represents the difference in amplification cycles required to detect the *WISP* genes in peripheral blood lymphocyte DNA compared with colon tumor DNA or colon tumor RNA compared with normal mucosal RNA. The Δ -method was used for calculation of the SE of the gene copy number or RNA expression level. The *WISP*-specific signal was normalized to that of the glyceraldehyde-3-phosphate dehydrogenase housekeeping gene. All TaqMan assay reagents were obtained from Perkin-Elmer Applied Biosystems.

RESULTS

Isolation of *WISP-1* and *WISP-2* by SSH. To identify Wnt-1-inducible genes, we used the technique of SSH using the

mouse mammary epithelial cell line C57MG and C57MG cells that stably express Wnt-1 (11). Candidate differentially expressed cDNAs (1,384 total) were sequenced. Thirty-nine percent of the sequences matched known genes or homologues, 32% matched expressed sequence tags, and 29% had no match. To confirm that the transcript was differentially expressed, semiquantitative reverse transcription-PCR and Northern analysis were performed by using mRNA from the C57MG and C57MG/Wnt-1 cells.

Two of the cDNAs, *WISP-1* and *WISP-2*, were differentially expressed, being induced in the C57MG/Wnt-1 cell line, but not in the parent C57MG cells or C57MG cells overexpressing Wnt-4 (Fig. 1A and B). Wnt-4, unlike Wnt-1, does not induce the morphological transformation of C57MG cells and has no effect on β -catenin levels (13, 14). Expression of *WISP-1* was up-regulated approximately 3-fold in the C57MG/Wnt-1 cell line and *WISP-2* by approximately 5-fold by both Northern analysis and reverse transcription-PCR.

An independent, but similar, system was used to examine *WISP* expression after Wnt-1 induction. C57MG cells expressing the *Wnt-1* gene under the control of a tetracycline-repressible promoter produce low amounts of Wnt-1 in the repressed state but show a strong induction of *Wnt-1* mRNA and protein within 24 hr after tetracycline removal (8). The levels of Wnt-1 and *WISP* RNA isolated from these cells at various times after tetracycline removal were assessed by quantitative PCR. Strong induction of Wnt-1 mRNA was seen as early as 10 hr after tetracycline removal. Induction of *WISP* mRNA (2- to 6-fold) was seen at 48 and 72 hr (data not shown). These data support our previous observations that show that *WISP* induction is correlated with Wnt-1 expression. Because the induction is slow, occurring after approximately 48 hr, the induction of *WISPs* may be an indirect response to Wnt-1 signaling.

cDNA clones of human *WISP-1* were isolated and the sequence compared with mouse *WISP-1*. The cDNA sequences of mouse and human *WISP-1* were 1,766 and 2,830 bp in length, respectively, and encode proteins of 367 aa, with predicted relative molecular masses of $\approx 40,000$ (M_r 40 K). Both have hydrophobic N-terminal signal sequences, 38 conserved cysteine residues, and four potential N-linked glycosylation sites and are 84% identical (Fig. 2A).

Full-length cDNA clones of mouse and human *WISP-2* were 1,734 and 1,293 bp in length, respectively, and encode proteins of 251 and 250 aa, respectively, with predicted relative molecular masses of $\approx 27,000$ (M_r 27 K) (Fig. 2B). Mouse and human *WISP-2* are 73% identical. Human *WISP-2* has no potential N-linked glycosylation sites, and mouse *WISP-2* has one at

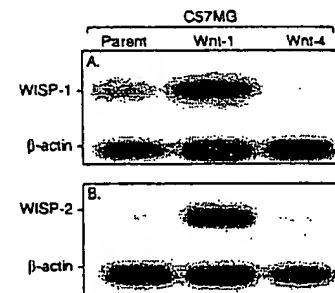


FIG. 1. *WISP-1* and *WISP-2* are induced by Wnt-1, but not Wnt-4, expression in C57MG cells. Northern analysis of *WISP-1* (A) and *WISP-2* (B) expression in C57MG, C57MG/Wnt-1, and C57MG/Wnt-4 cells. Poly(A)⁺ RNA (2 μ g) was subjected to Northern blot analysis and hybridized with a 70-bp mouse *WISP-1*-specific probe (amino acids 278–300) or a 190-bp *WISP-2*-specific probe (nucleotides 1438–1627) in the 3' untranslated region. Blots were rehybridized with human β -actin probe.

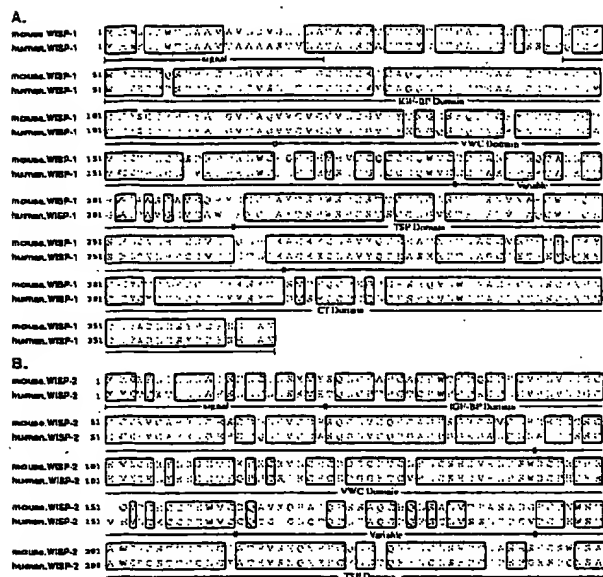


FIG. 2. Encoded amino acid sequence alignment of mouse and human *WISP-1* (A) and mouse and human *WISP-2* (B). The potential signal sequence, insulin-like growth factor-binding protein (IGF-BP), VWC, thrombospondin (TSP), and C-terminal (CT) domains are underlined.

position 197. *WISP-2* has 28 cysteine residues that are conserved among the 38 cysteines found in *WISP-1*.

Identification of *WISP-3*. To search for related proteins, we screened expressed sequence tag (EST) databases with the *WISP-1* protein sequence and identified several ESTs as potentially related sequences. We identified a homologous protein that we have called *WISP-3*. A full-length human *WISP-3* cDNA of 1,371 bp was isolated corresponding to those ESTs that encode a 354-aa protein with a predicted molecular mass of 39,293. *WISP-3* has two potential N-linked glycosylation sites and 36 cysteine residues. An alignment of the three human *WISP* proteins shows that *WISP-1* and *WISP-3* are the most similar (42% identity), whereas *WISP-2* has 37% identity with *WISP-1* and 32% identity with *WISP-3* (Fig. 3A).

***WISPs* Are Homologous to the CTGF Family of Proteins.** Human *WISP-1*, *WISP-2*, and *WISP-3* are novel sequences; however, mouse *WISP-1* is the same as the recently identified *Elm1* gene. *Elm1* is expressed in low, but not high, metastatic mouse melanoma cells, and suppresses the *in vivo* growth and metastatic potential of K-1735 mouse melanoma cells (15). Human and mouse *WISP-2* are homologous to the recently described rat gene, *rCop-1* (16). Significant homology (36–44%) was seen to the CCN family of growth factors. This family includes three members, CTGF, Cyr61, and the protooncogene *nov*. CTGF is a chemotactic and mitogenic factor for fibroblasts that is implicated in wound healing and fibrotic disorders and is induced by TGF- β (17). Cyr61 is an extracellular matrix signaling molecule that promotes cell adhesion, proliferation, migration, angiogenesis, and tumor growth (18, 19). *nov* (nephroblastoma overexpressed) is an immediate early gene associated with quiescence and found altered in Wilms tumors (20). The proteins of the CCN family share functional, but not sequence, similarity to Wnt-1. All are secreted, cysteine-rich heparin binding glycoproteins that associate with the cell surface and extracellular matrix.

WISP proteins exhibit the modular architecture of the CCN family, characterized by four conserved cysteine-rich domains (Fig. 3B) (21). The N-terminal domain, which includes the first 12 cysteine residues, contains a consensus sequence (GCGC-CXXC) conserved in most insulin-like growth factor (IGF)-

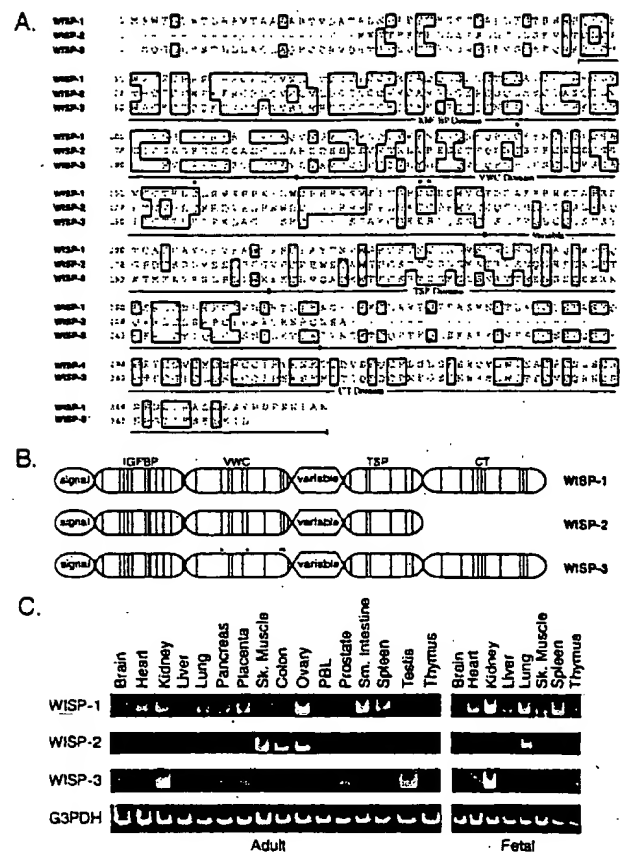


FIG. 3. (A) Encoded amino acid sequence alignment of human *WISPs*. The cysteine residues of *WISP-1* and *WISP-2* that are not present in *WISP-3* are indicated with a dot. (B) Schematic representation of the *WISP* proteins showing the domain structure and cysteine residues (vertical lines). The four cysteine residues in the VWC domain that are absent in *WISP-3* are indicated with a dot. (C) Expression of *WISP* mRNA in human tissues. PCR was performed on human multiple-tissue cDNA panels (CLONTECH) from the indicated adult and fetal tissues.

binding proteins (BP). This sequence is conserved in *WISP-2* and *WISP-3*, whereas *WISP-1* has a glutamine in the third position instead of a glycine. CTGF recently has been shown to specifically bind IGF (22) and a truncated *nov* protein lacking the IGF-BP domain is oncogenic (23). The von Willebrand factor type C module (VWC), also found in certain collagens and mucins, covers the next 10 cysteine residues, and is thought to participate in protein complex formation and oligomerization (24). The VWC domain of *WISP-3* differs from all CCN family members described previously, in that it contains only six of the 10 cysteine residues (Fig. 3A and B). A short variable region follows the VWC domain. The third module, the thrombospondin (TSP) domain is involved in binding to sulfated glycoconjugates and contains six cysteine residues and a conserved WSxCSxCG motif first identified in thrombospondin (25). The C-terminal (CT) module containing the remaining 10 cysteines is thought to be involved in dimerization and receptor binding (26). The CT domain is present in all CCN family members described to date but is absent in *WISP-2* (Fig. 3A and B). The existence of a putative signal sequence and the absence of a transmembrane domain suggest that *WISPs* are secreted proteins, an observation supported by an analysis of their expression and secretion from mammalian cell and baculovirus cultures (data not shown).

Expression of *WISP* mRNA in Human Tissues. Tissue-specific expression of human *WISPs* was characterized by PCR

analysis on adult and fetal multiple tissue cDNA panels. *WISP-1* expression was seen in the adult heart, kidney, lung, pancreas, placenta, ovary, small intestine, and spleen (Fig. 3C). Little or no expression was detected in the brain, liver, skeletal muscle, colon, peripheral blood leukocytes, prostate, testis, or thymus. *WISP-2* had a more restricted tissue expression and was detected in adult skeletal muscle, colon, ovary, and fetal lung. Predominant expression of *WISP-3* was seen in adult kidney and testis and fetal kidney. Lower levels of *WISP-3* expression were detected in placenta, ovary, prostate, and small intestine.

In Situ Localization of *WISP-1* and *WISP-2*. Expression of *WISP-1* and *WISP-2* was assessed by *in situ* hybridization in mammary tumors from Wnt-1 transgenic mice. Strong expression of *WISP-1* was observed in stromal fibroblasts lying within the fibrovascular tumor stroma (Fig. 4A–D). However, low-level *WISP-1* expression also was observed focally within tumor cells (data not shown). No expression was observed in normal breast. Like *WISP-1*, *WISP-2* expression also was seen in the tumor stroma in breast tumors from Wnt-1 transgenic animals (Fig. 4E–H). However, *WISP-2* expression in the stroma was in spindle-shaped cells adjacent to capillary vessels, whereas

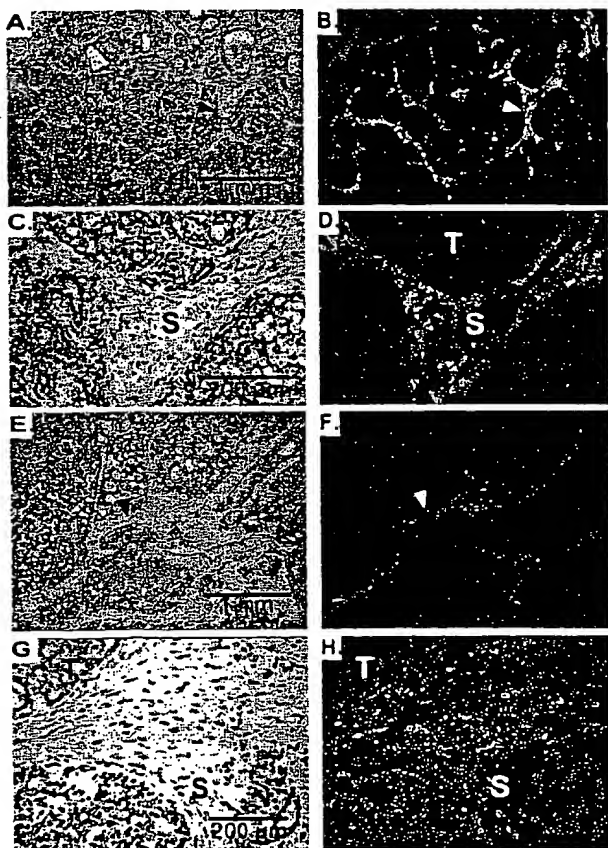


FIG. 4. (A, C, E, and G) Representative hematoxylin/eosin-stained images from breast tumors in Wnt-1 transgenic mice. The corresponding dark-field images showing *WISP-1* expression are shown in B and D. The tumor is a moderately well-differentiated adenocarcinoma showing evidence of adenoid cystic change. At low power (A and B), expression of *WISP-1* is seen in the delicate branching fibrovascular tumor stroma (arrowhead). At higher magnification, expression is seen in the stromal(s) fibroblasts (C and D), and tumor cells are negative. Focal expression of *WISP-1*, however, was observed in tumor cells in some areas. Images of *WISP-2* expression are shown in E–H. At low power (E and F), expression of *WISP-2* is seen in cells lying within the fibrovascular tumor stroma. At higher magnification, these cells appeared to be adjacent to capillary vessels whereas tumor cells are negative (G and H).

the predominant cell type expressing *WISP-1* was the stromal fibroblasts.

Chromosome Localization of the *WISP* Genes. The chromosomal location of the human *WISP* genes was determined by radiation hybrid mapping panels. *WISP-1* is approximately 3.48 cR from the meiotic marker AFM259xc5 [logarithm of odds (lod) score 16.31] on chromosome 8q24.1 to 8q24.3, in the same region as the human locus of the *novH* family member (27) and roughly 4 Mbs distal to *c-myc* (28). Preliminary fine mapping indicates that *WISP-1* is located near D8S1712 STS. *WISP-2* is linked to the marker SHGC-33922 (lod = 1,000) on chromosome 20q12–20q13.1. Human *WISP-3* mapped to chromosome 6q22–6q23 and is linked to the marker AFM211ze5 (lod = 1,000). *WISP-3* is approximately 18 Mbs proximal to CTGF and 23 Mbs proximal to the human cellular oncogene *MYB* (27, 29).

Amplification and Aberrant Expression of *WISPs* in Human Colon Tumors. Amplification of protooncogenes is seen in many human tumors and has etiological and prognostic significance. For example, in a variety of tumor types, *c-myc* amplification has been associated with malignant progression and poor prognosis (30). Because *WISP-1* resides in the same general chromosomal location (8q24) as *c-myc*, we asked whether it was a target of gene amplification, and, if so, whether this amplification was independent of the *c-myc* locus. Genomic DNA from human colon cancer cell lines was assessed by quantitative PCR and Southern blot analysis. (Fig. 5A and B). Both methods detected similar degrees of *WISP-1* amplification. Most cell lines showed significant (2- to 4-fold) amplification, with the HT-29 and WiDr cell lines demonstrating an 8-fold increase. Significantly, the pattern of amplification observed did not correlate with that observed for *c-myc*, indicating that the *c-myc* gene is not part of the amplicon that involves the *WISP-1* locus.

We next examined whether the *WISP* genes were amplified in a panel of 25 primary human colon adenocarcinomas. The relative *WISP* gene copy number in each colon tumor DNA was compared with pooled normal DNA from 10 donors by quantitative PCR (Fig. 6). The copy number of *WISP-1* and *WISP-2* was significantly greater than one, approximately 2-fold for *WISP-1* in about 60% of the tumors and 2- to 4-fold for *WISP-2* in 92% of the tumors ($P < 0.001$ for each). The copy number for *WISP-3* was indistinguishable from one ($P = 0.166$). In addition, the copy number of *WISP-2* was significantly higher than that of *WISP-1* ($P < 0.001$).

The levels of *WISP* transcripts in RNA isolated from 19 adenocarcinomas and their matched normal mucosa were

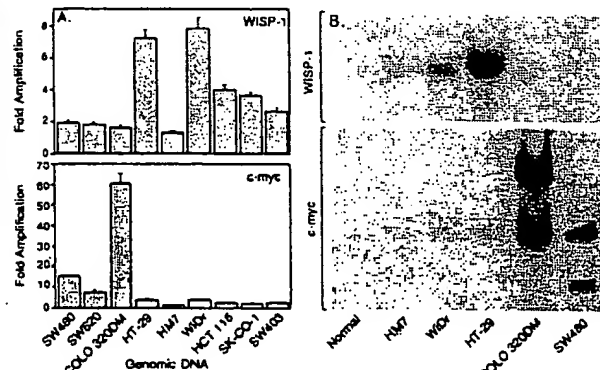


FIG. 5. Amplification of *WISP-1* genomic DNA in colon cancer cell lines. (A) Amplification in cell line DNA was determined by quantitative PCR. (B) Southern blots containing genomic DNA (10 μ g) digested with *Eco*RI (*WISP-1*) or *Xba*I (*c-myc*) were hybridized with a 100-bp human *WISP-1* probe (amino acids 186–219) or a human *c-myc* probe (located at bp 1901–2000). The *WISP* and *myc* genes are detected in normal human genomic DNA after a longer film exposure.

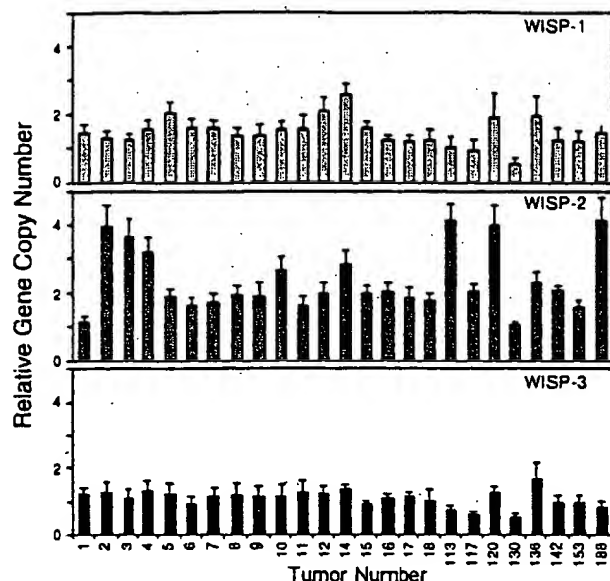


FIG. 6. Genomic amplification of *WISP* genes in human colon tumors. The relative gene copy number of the *WISP* genes in 25 adenocarcinomas was assayed by quantitative PCR, by comparing DNA from primary human tumors with pooled DNA from 10 healthy donors. The data are means \pm SEM from one experiment done in triplicate. The experiment was repeated at least three times.

assessed by quantitative PCR (Fig. 7). The level of *WISP-1* RNA present in tumor tissue varied but was significantly increased (2- to >25-fold) in 84% (16/19) of the human colon tumors examined compared with normal adjacent mucosa. Four of 19 tumors showed greater than 10-fold overexpression. In contrast, in 79% (15/19) of the tumors examined, *WISP-2* RNA expression was significantly lower in the tumor than the mucosa. Similar to *WISP-1*, *WISP-3* RNA was overexpressed in 63% (12/19) of the colon tumors compared with the normal

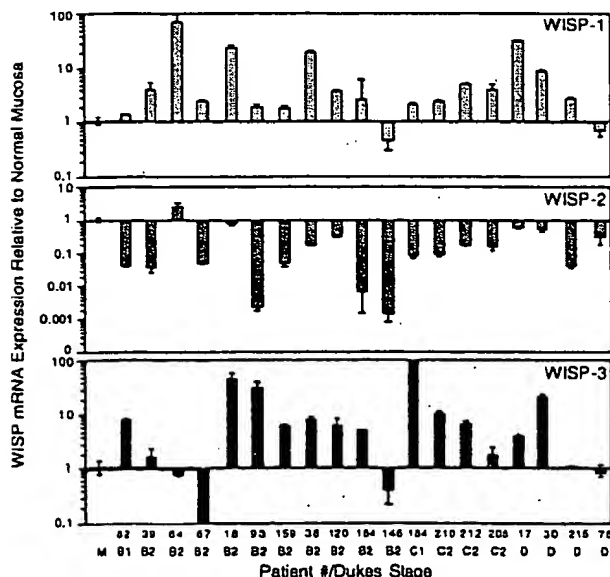


FIG. 7. *WISP* RNA expression in primary human colon tumors relative to expression in normal mucosa from the same patient. Expression of *WISP* mRNA in 19 adenocarcinomas was assayed by quantitative PCR. The Dukes stage of the tumor is listed under the sample number. The data are means \pm SEM from one experiment done in triplicate. The experiment was repeated at least twice.

mucosa. The amount of overexpression of *WISP-3* ranged from 4- to >40-fold.

DISCUSSION

One approach to understanding the molecular basis of cancer is to identify differences in gene expression between cancer cells and normal cells. Strategies based on assumptions that steady-state mRNA levels will differ between normal and malignant cells have been used to clone differentially expressed genes (31). We have used a PCR-based selection strategy, SSH, to identify genes selectively expressed in C57MG mouse mammary epithelial cells transformed by Wnt-1.

Three of the genes isolated, *WISP-1*, *WISP-2*, and *WISP-3*, are members of the CCN family of growth factors, which includes CTGF, Cyr61, and *nov*, a family not previously linked to Wnt signaling.

Two independent experimental systems demonstrated that *WISP* induction was associated with the expression of Wnt-1. The first was C57MG cells infected with a Wnt-1 retroviral vector or C57MG cells expressing Wnt-1 under the control of a tetracycline-repressible promoter, and the second was in Wnt-1 transgenic mice, where breast tissue expresses Wnt-1, whereas normal breast tissue does not. No *WISP* RNA expression was detected in mammary tumors induced by polyoma virus middle T antigen (data not shown). These data suggest a link between Wnt-1 and *WISPs* in that in these two situations, *WISP* induction was correlated with Wnt-1 expression.

It is not clear whether the *WISPs* are directly or indirectly induced by the downstream components of the Wnt-1 signaling pathway (i.e., β -catenin-TCF-1/Lef1). The increased levels of *WISP* RNA were measured in Wnt-1-transformed cells, hours or days after Wnt-1 transformation. Thus, *WISP* expression could result from Wnt-1 signaling directly through β -catenin transcription factor regulation or alternatively through Wnt-1 signaling turning on a transcription factor, which in turn regulates *WISPs*.

The *WISPs* define an additional subfamily of the CCN family of growth factors. One striking difference observed in the protein sequence of *WISP-2* is the absence of a CT domain, which is present in CTGF, Cyr61, *nov*, *WISP-1*, and *WISP-3*. This domain is thought to be involved in receptor binding and dimerization. Growth factors, such as TGF- β , platelet-derived growth factor, and nerve growth factor, which contain a cystine knot motif exist as dimers (32). It is tempting to speculate that *WISP-1* and *WISP-3* may exist as dimers, whereas *WISP-2* exists as a monomer. If the CT domain is also important for receptor binding, *WISP-2* may bind its receptor through a different region of the molecule than the other CCN family members. No specific receptors have been identified for CTGF or *nov*. A recent report has shown that integrin $\alpha\beta_3$ serves as an adhesion receptor for Cyr61 (33).

The strong expression of *WISP-1* and *WISP-2* in cells lying within the fibrovascular tumor stroma in breast tumors from Wnt-1 transgenic animals is consistent with previous observations that transcripts for the related CTGF gene are primarily expressed in the fibrous stroma of mammary tumors (34). Epithelial cells are thought to control the proliferation of connective tissue stroma in mammary tumors by a cascade of growth factor signals similar to that controlling connective tissue formation during wound repair. It has been proposed that mammary tumor cells or inflammatory cells at the tumor interstitial interface secrete TGF- β 1, which is the stimulus for stromal proliferation (34). TGF- β 1 is secreted by a large percentage of malignant breast tumors and may be one of the growth factors that stimulates the production of CTGF and *WISPs* in the stroma.

It was of interest that *WISP-1* and *WISP-2* expression was observed in the stromal cells that surrounded the tumor cells

(epithelial cells) in the Wnt-1 transgenic mouse sections of breast tissue. This finding suggests that paracrine signaling could occur in which the stromal cells could supply WISP-1 and WISP-2 to regulate tumor cell growth on the WISP extracellular matrix. Stromal cell-derived factors in the extracellular matrix have been postulated to play a role in tumor cell migration and proliferation (35). The localization of *WISP-1* and *WISP-2* in the stromal cells of breast tumors supports this paracrine model.

An analysis of *WISP-1* gene amplification and expression in human colon tumors showed a correlation between DNA amplification and overexpression; whereas overexpression of *WISP-3* RNA was seen in the absence of DNA amplification. In contrast, *WISP-2* DNA was amplified in the colon tumors, but its mRNA expression was significantly reduced in the majority of tumors compared with the expression in normal colonic mucosa from the same patient. The gene for human *WISP-2* was localized to chromosome 20q12-20q13, at a region frequently amplified and associated with poor prognosis in node negative breast cancer and many colon cancers, suggesting the existence of one or more oncogenes at this locus (36-38). Because the center of the 20q13 amplicon has not yet been identified, it is possible that the apparent amplification observed for *WISP-2* may be caused by another gene in this amplicon.

A recent manuscript on *rCop-1*, the rat orthologue of *WISP-2*, describes the loss of expression of this gene after cell transformation, suggesting it may be a negative regulator of growth in cell lines (16). Although the mechanism by which *WISP-2* RNA expression is down-regulated during malignant transformation is unknown, the reduced expression of *WISP-2* in colon tumors and cell lines suggests that it may function as a tumor suppressor. These results show that the *WISP* genes are aberrantly expressed in colon cancer and suggest that their altered expression may confer selective growth advantage to the tumor.

Members of the Wnt signaling pathway have been implicated in the pathogenesis of colon cancer, breast cancer, and melanoma, including the tumor suppressor gene adenomatous polyposis coli and β -catenin (39). Mutations in specific regions of either gene can cause the stabilization and accumulation of cytoplasmic β -catenin, which presumably contributes to human carcinogenesis through the activation of target genes such as the *WISPs*. Although the mechanism by which Wnt-1 transforms cells and induces tumorigenesis is unknown, the identification of *WISPs* as genes that may be regulated downstream of Wnt-1 in C57MG cells suggests they could be important mediators of Wnt-1 transformation. The amplification and altered expression patterns of the *WISPs* in human colon tumors may indicate an important role for these genes in tumor development.

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methods. Peptides AENK or AEQK were dissolved in water, made isotonic with NaCl and diluted into RPMI growth medium. T-cell-proliferation assays were done essentially as described^{20,21}. Briefly, after antigen pulsing ($30 \mu\text{g ml}^{-1}$ TTCF) with tetrapeptides ($1\text{--}2 \text{ mg ml}^{-1}$), PBMCs or EBV-B cells were washed in PBS and fixed for 45 s in 0.05% glutaraldehyde. Glycine was added to a final concentration of 0.1M and the cells were washed five times in RPMI 1640 medium containing 1% FCS before co-culture with T-cell clones in round-bottom 96-well microtitre plates. After 48 h, the cultures were pulsed with $1 \mu\text{Ci}$ of ^3H -thymidine and harvested for scintillation counting 16 h later. Predigestion of native TTCF was done by incubating $200 \mu\text{g}$ TTCF with $0.25 \mu\text{g}$ pig kidney legumain in $500 \mu\text{l}$ 50 mM citrate buffer, pH 5.5, for 1 h at 37°C . **Glycopeptide digestions.** The peptides HIDNEEDI, HIDN(N-glucosamine) EEDI and HIDNESDI, which are based on the TTCF sequence, and QQQLFGSNVTDCSGNFCLFR(KKK), which is based on human transferrin, were obtained by custom synthesis. The three C-terminal lysine residues were added to the natural sequence to aid solubility. The transferrin glycopeptide QQQLFGSNVTDCSGNFCLFR was prepared by tryptic (Promega) digestion of 5 mg reduced, carboxy-methylated human transferrin followed by concanavalin A chromatography¹¹. Glycopeptides corresponding to residues 622–642 and 421–452 were isolated by reverse-phase HPLC and identified by mass spectrometry and N-terminal sequencing. The lyophilized transferrin-derived peptides were redissolved in 50 mM sodium acetate, pH 5.5, 10 mM dithiothreitol, 20% methanol. Digestions were performed for 3 h at 30°C with $5\text{--}50 \text{ mU ml}^{-1}$ pig kidney legumain or B-cell AEP. Products were analysed by HPLC or MALDI-TOF mass spectrometry using a matrix of 10 mg ml^{-1} α -cyanocinnamic acid in 50% acetonitrile/0.1% TFA and a PerSeptive Biosystems Elite STR mass spectrometer set to linear or reflector mode. Internal standardization was obtained with a matrix ion of 568.13 mass units.

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Genomic amplification of a decoy receptor for Fas ligand in lung and colon cancer

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Fas ligand (FasL) is produced by activated T cells and natural killer cells and it induces apoptosis (programmed cell death) in target cells through the death receptor Fas/Apo1/CD95 (ref. 1). One important role of FasL and Fas is to mediate immune-cytotoxic killing of cells that are potentially harmful to the organism, such as virus-infected or tumour cells¹. Here we report the discovery of a soluble decoy receptor, termed decoy receptor 3 (Dcr3), that binds to FasL and inhibits FasL-induced apoptosis. The Dcr3 gene was amplified in about half of 35 primary lung and colon tumours studied, and Dcr3 messenger RNA was expressed in malignant tissue. Thus, certain tumours may escape FasL-dependent immune-cytotoxic attack by expressing a decoy receptor that blocks FasL.

By searching expressed sequence tag (EST) databases, we identified a set of related ESTs that showed homology to the tumour necrosis factor (TNF) receptor (TNFR) gene superfamily². Using the overlapping sequence, we isolated a previously unknown full-length complementary DNA from human fetal lung. We named the protein encoded by this cDNA decoy receptor 3 (Dcr3). The cDNA encodes a 300-amino-acid polypeptide that resembles members of the TNFR family (Fig. 1a): the amino terminus contains a leader sequence, which is followed by four tandem cysteine-rich domains (CRDs). Like one other TNFR homologue, osteoprotegerin (OPG)³, Dcr3 lacks an apparent transmembrane sequence, which indicates that it may be a secreted, rather than a membrane-associated, molecule. We expressed a recombinant, histidine-tagged form of Dcr3 in mammalian cells; Dcr3 was secreted into the cell culture medium, and migrated on polyacrylamide gels as a protein of relative molecular mass 35,000 (data not shown). Dcr3 shares sequence identity in particular with OPG (31%) and TNFR2 (29%), and has relatively less homology with Fas (17%). All of the cysteines in the four CRDs of Dcr3 and OPG are conserved; however, the carboxy-terminal portion of Dcr3 is 101 residues shorter.

We analysed expression of Dcr3 mRNA in human tissues by northern blotting (Fig. 1b). We detected a predominant 1.2-kilobase transcript in fetal lung, brain, and liver, and in adult spleen, colon and lung. In addition, we observed relatively high Dcr3 mRNA expression in the human colon carcinoma cell line SW480.

To investigate potential ligand interactions of Dcr3, we generated a recombinant, Fc-tagged Dcr3 protein. We tested binding of Dcr3-Fc to human 293 cells transfected with individual TNF-family ligands, which are expressed as type 2 transmembrane proteins (these transmembrane proteins have their N termini in the cytosol). Dcr3-Fc showed a significant increase in binding to cells transfected with FasL⁴ (Fig. 2a), but not to cells transfected with TNF⁵, Apo2L/TRAIL^{6,7}, Apo3L/TWEAK^{8,9}, or OPGL/TRACE/

RANKL¹⁰⁻¹² (data not shown). DcR3-Fc immunoprecipitated shed FasL from FasL-transfected 293 cells (Fig. 2b) and purified soluble FasL (Fig. 2c), as did the Fc-tagged ectodomain of Fas but not TNFR1. Gel-filtration chromatography showed that DcR3-Fc and soluble FasL formed a stable complex (Fig. 2d). Equilibrium analysis indicated that DcR3-Fc and Fas-Fc bound to soluble FasL with a comparable affinity ($K_d = 0.8 \pm 0.2$ and 1.1 ± 0.1 nM, respectively; Fig. 2e), and that DcR3-Fc could block nearly all of the binding of soluble FasL to Fas-Fc (Fig. 2e, inset). Thus, DcR3 competes with Fas for binding to FasL.

To determine whether binding of DcR3 inhibits FasL activity, we tested the effect of DcR3-Fc on apoptosis induction by soluble FasL in Jurkat T leukaemia cells, which express Fas (Fig. 3a). DcR3-Fc and Fas-Fc blocked soluble-FasL-induced apoptosis in a similar dose-dependent manner, with half-maximal inhibition at $\sim 0.1 \mu\text{g ml}^{-1}$. Time-course analysis showed that the inhibition did not merely delay cell death, but rather persisted for at least 24 hours (Fig. 3b). We also tested the effect of DcR3-Fc on activation-induced cell death (AICD) of mature T lymphocytes, a FasL-dependent process¹. Consistent with previous results¹³, activation of interleukin-2-stimulated CD4-positive T cells with anti-CD3 antibody increased the level of apoptosis twofold, and Fas-Fc blocked this effect substantially (Fig. 3c); DcR3-Fc blocked the

induction of apoptosis to a similar extent. Thus, DcR3 binding blocks apoptosis induction by FasL.

FasL-induced apoptosis is important in elimination of virus-infected cells and cancer cells by natural killer cells and cytotoxic T lymphocytes; an alternative mechanism involves perforin and granzymes¹⁴⁻¹⁶. Peripheral blood natural killer cells triggered marked cell death in Jurkat T leukaemia cells (Fig. 3d); DcR3-Fc and Fas-Fc each reduced killing of target cells from $\sim 65\%$ to $\sim 30\%$, with half-maximal inhibition at $\sim 1 \mu\text{g ml}^{-1}$; the residual killing was probably mediated by the perforin/granzyme pathway. Thus, DcR3 binding blocks FasL-dependent natural killer cell activity. Higher DcR3-Fc and Fas-Fc concentrations were required to block natural killer cell activity compared with those required to block soluble FasL activity, which is consistent with the greater potency of membrane-associated FasL compared with soluble FasL¹⁷.

Given the role of immune-cytotoxic cells in elimination of tumour cells and the fact that DcR3 can act as an inhibitor of FasL, we proposed that DcR3 expression might contribute to the ability of some tumours to escape immune-cytotoxic attack. As genomic amplification frequently contributes to tumorigenesis, we investigated whether the DcR3 gene is amplified in cancer. We analysed DcR3 gene-copy number by quantitative polymerase chain

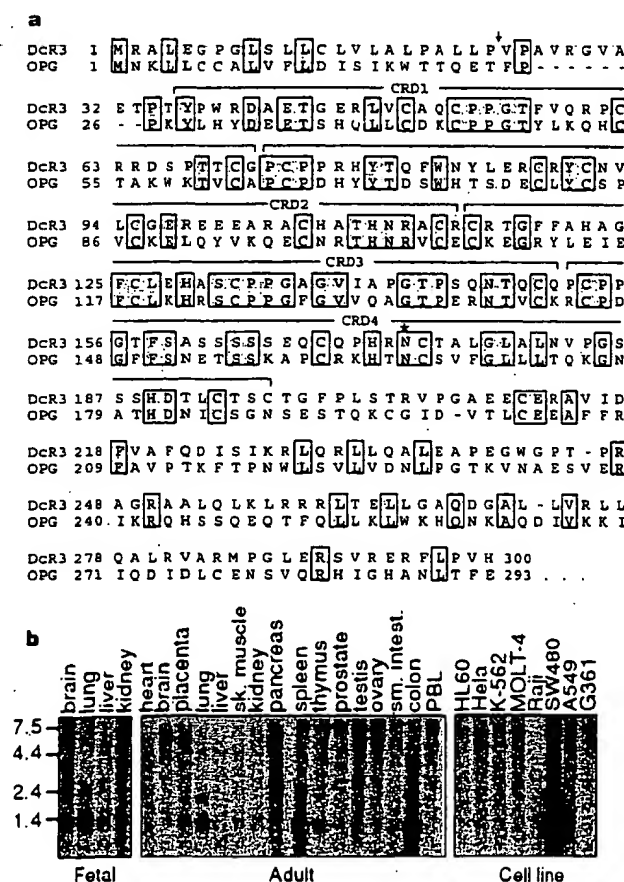


Figure 1 Primary structure and expression of human DcR3. **a**, Alignment of the amino-acid sequences of DcR3 and of osteoprotegerin (OPG); the C-terminal 101 residues of OPG are not shown. The putative signal cleavage site (arrow), the cysteine-rich domains (CRD 1-4), and the *N*-linked glycosylation site (asterisk) are shown. **b**, Expression of DcR3 mRNA. Northern hybridization analysis was done using the DcR3 cDNA as a probe and blots of poly(A)⁺ RNA (Clontech) from human fetal and adult tissues or cancer cell lines. PBL, peripheral blood lymphocyte.

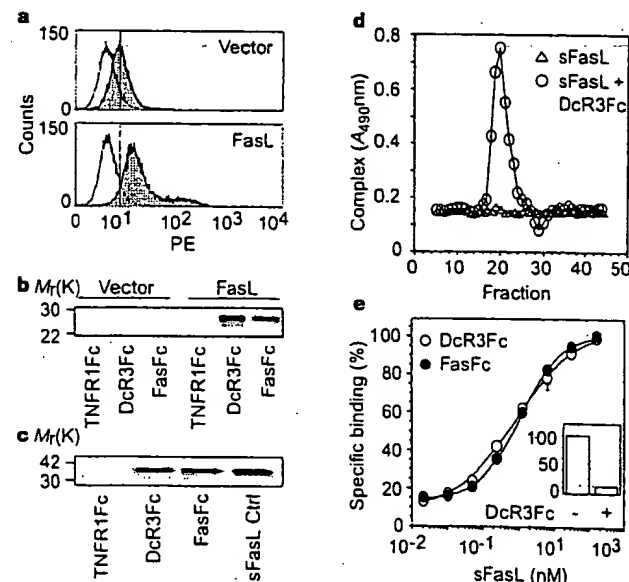


Figure 2 Interaction of DcR3 with FasL. **a**, 293 cells were transfected with pRK5 vector (top) or with pRK5 encoding full-length FasL (bottom), incubated with DcR3-Fc (solid line, shaded area), TNFR1-Fc (dotted line) or buffer control (dashed line) (the dashed and dotted lines overlap), and analysed for binding by FACS. Statistical analysis showed a significant difference ($P < 0.001$) between the binding of DcR3-Fc to cells transfected with FasL or pRK5. PE, phycoerythrin-labelled cells. **b**, 293 cells were transfected as in **a** and metabolically labelled, and cell supernatants were immunoprecipitated with Fc-tagged TNFR1, DcR3 or Fas. **c**, Purified soluble FasL (sFasL) was immunoprecipitated with TNFR1-Fc, DcR3-Fc or Fas-Fc and visualized by immunoblot with anti-FasL antibody. sFasL was loaded directly for comparison in the right-hand lane. **d**, Flag-tagged sFasL was incubated with DcR3-Fc or with buffer and resolved by gel filtration; column fractions were analysed in an assay that detects complexes containing DcR3-Fc and sFasL-Flag. **e**, Equilibrium binding of DcR3-Fc or Fas-Fc to sFasL-Flag. Inset, competition of DcR3-Fc with Fas-Fc for binding to sFasL-Flag.

reaction (PCR)¹⁸ in genomic DNA from 35 primary lung and colon tumours, relative to pooled genomic DNA from peripheral blood leukocytes (PBLs) of 10 healthy donors. Eight of 18 lung tumours and 9 of 17 colon tumours showed DcR3 gene amplification, ranging from 2- to 18-fold (Fig. 4a, b). To confirm this result, we analysed the colon tumour DNAs with three more, independent sets of DcR3-based PCR primers and probes; we observed nearly the same amplification (data not shown).

We then analysed DcR3 mRNA expression in primary tumour tissue sections by *in situ* hybridization. We detected DcR3 expression in 6 out of 15 lung tumours, 2 out of 2 colon tumours, 2 out of 5 breast tumours, and 1 out of 1 gastric tumour (data not shown). A section through a squamous-cell carcinoma of the lung is shown in Fig. 4c. DcR3 mRNA was localized to infiltrating malignant epithelium, but was essentially absent from adjacent stroma, indicating tumour-specific expression. Although the individual tumour specimens that we analysed for mRNA expression and gene amplification were different, the *in situ* hybridization results are consistent with the finding that the DcR3 gene is amplified frequently in tumours. SW480 colon carcinoma cells, which showed abundant DcR3 mRNA expression (Fig. 1b), also had marked DcR3 gene amplification, as shown by quantitative PCR (fourfold) and by Southern blot hybridization (fivefold) (data not shown).

If DcR3 amplification in cancer is functionally relevant, then DcR3 should be amplified more than neighbouring genomic regions that are not important for tumour survival. To test this,

we mapped the human DcR3 gene by radiation-hybrid analysis; DcR3 showed linkage to marker AFM218xe7 (T160), which maps to chromosome position 20q13. Next, we isolated from a bacterial artificial chromosome (BAC) library a human genomic clone that carries DcR3, and sequenced the ends of the clone's insert. We then determined, from the nine colon tumours that showed twofold or greater amplification of DcR3, the copy number of the DcR3-flanking sequences (reverse and forward) from the BAC, and of seven genomic markers that span chromosome 20 (Fig. 4d). The DcR3-linked reverse marker showed an average amplification of roughly threefold, slightly less than the approximately fourfold amplification of DcR3; the other markers showed little or no amplification. These data indicate that DcR3 may be at the 'epicentre' of a distal chromosome 20 region that is amplified in colon cancer, consistent with the possibility that DcR3 amplification promotes tumour survival.

Our results show that DcR3 binds specifically to FasL and inhibits FasL activity. We did not detect DcR3 binding to several other TNF-ligand-family members; however, this does not rule out the possibility that DcR3 interacts with other ligands, as do some other TNFR family members, including OPG^{2,19}.

FasL is important in regulating the immune response; however, little is known about how FasL function is controlled. One mechanism involves the molecule cFLIP, which modulates apoptosis signalling downstream of Fas²⁰. A second mechanism involves proteolytic shedding of FasL from the cell surface¹⁷. DcR3 competes with Fas for

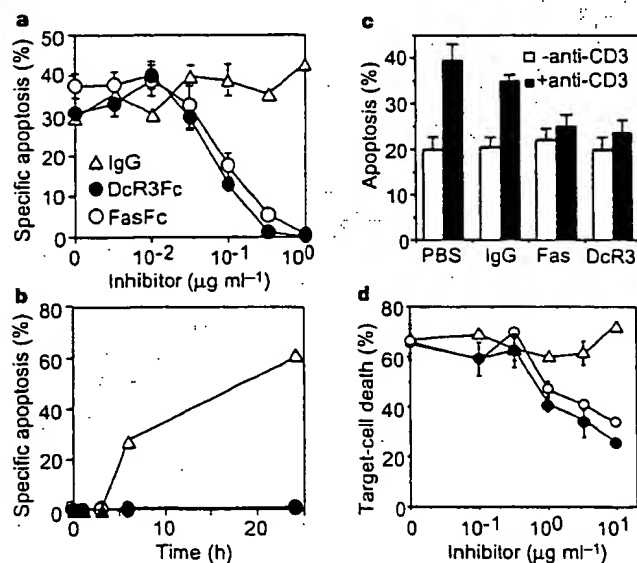


Figure 3 Inhibition of FasL activity by DcR3. **a**, Human Jurkat T leukaemia cells were incubated with Flag-tagged soluble FasL (sFasL: 5 ng ml⁻¹) oligomerized with anti-Flag antibody (0.1 μg ml⁻¹) in the presence of the proposed inhibitors DcR3-Fc, Fas-Fc or human IgG1 and assayed for apoptosis (mean ± s.e.m. of triplicates). **b**, Jurkat cells were incubated with sFasL-Flag plus anti-Flag antibody as in **a**, in presence of 1 μg ml⁻¹ DcR3-Fc (filled circles), Fas-Fc (open circles) or human IgG1 (triangles), and apoptosis was determined at the indicated time points. **c**, Peripheral blood T cells were stimulated with PHA and interleukin-2, followed by control (white bars) or anti-CD3 antibody (filled bars), together with phosphate-buffered saline (PBS), human IgG1, Fas-Fc, or DcR3-Fc (10 μg ml⁻¹). After 16 h, apoptosis of CD4⁺ cells was determined (mean ± s.e.m. of results from five donors). **d**, Peripheral blood natural killer cells were incubated with ⁵¹Cr-labelled Jurkat cells in the presence of DcR3-Fc (filled circles), Fas-Fc (open circles) or human IgG1 (triangles), and target-cell death was determined by release of ⁵¹Cr (mean ± s.d. for two donors, each in triplicate).

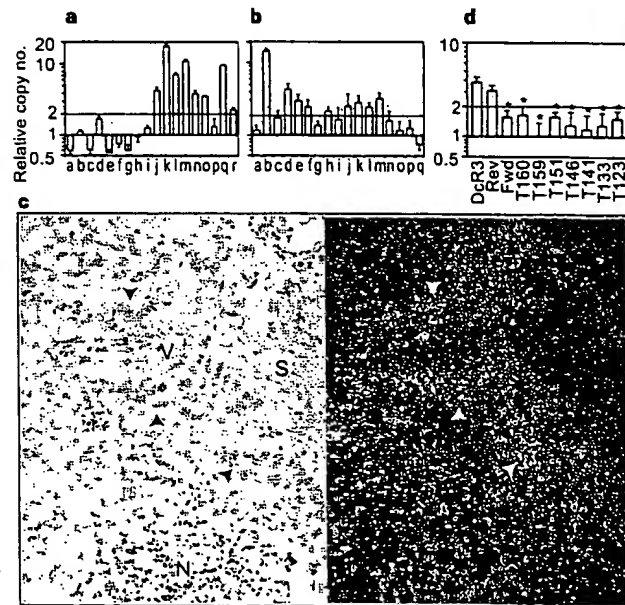


Figure 4 Genomic amplification of DcR3 in tumours. **a**, Lung cancers, comprising eight adenocarcinomas (c, d, f, g, h, j, k, r), seven squamous-cell carcinomas (a, e, m, n, o, p, q), one non-small-cell carcinoma (b), one small-cell carcinoma (i), and one bronchial adenocarcinoma (l). The data are means ± s.d. of 2 experiments done in duplicate. **b**, Colon tumours, comprising 17 adenocarcinomas. Data are means ± s.e.m. of five experiments done in duplicate. **c**, *In situ* hybridization analysis of DcR3 mRNA expression in a squamous-cell carcinoma of the lung. A representative bright-field image (left) and the corresponding dark-field image (right) show DcR3 mRNA over infiltrating malignant epithelium (arrowheads). Adjacent non-malignant stroma (S), blood vessel (V) and necrotic tumour tissue (N) are also shown. **d**, Average amplification of DcR3 compared with amplification of neighbouring genomic regions (reverse and forward, Rev and Fwd), the DcR3-linked marker T160, and other chromosome-20 markers, in the nine colon tumours showing DcR3 amplification of twofold or more (b). Data are from two experiments done in duplicate. Asterisk indicates $P < 0.01$ for a Student's *t*-test comparing each marker with DcR3.

FasL binding; hence, it may represent a third mechanism of extracellular regulation of FasL activity. A decoy receptor that modulates the function of the cytokine interleukin-1 has been described²¹. In addition, two decoy receptors that belong to the TNFR family, DcR1 and DcR2, regulate the FasL-related apoptosis-inducing molecule Apo2L²². Unlike DcR1 and DcR2, which are membrane-associated proteins, DcR3 is directly secreted into the extracellular space. One other secreted TNFR-family member is OPG², which shares greater sequence homology with DcR3 (31%) than do DcR1 (17%) or DcR2 (19%); OPG functions as a third decoy for Apo2L¹⁹. Thus, DcR3 and OPG define a new subset of TNFR-family members that function as secreted decoys to modulate ligands that induce apoptosis. Pox viruses produce soluble TNFR homologues that neutralize specific TNF-family ligands, thereby modulating the antiviral immune response². Our results indicate that a similar mechanism, namely, production of a soluble decoy receptor for FasL, may contribute to immune evasion by certain tumours. □

Methods

Isolation of DcR3 cDNA. Several overlapping ESTs in GenBank (accession numbers AA025672, AA025673 and W67560) and in LifeseqTM (Incyte Pharmaceuticals; accession numbers 1339238, 1533571, 1533650, 1542861, 1789372 and 2207027) showed similarity to members of the TNFR family. We screened human cDNA libraries by PCR with primers based on the region of EST consensus; fetal lung was positive for a product of the expected size. By hybridization to a PCR-generated probe based on the ESTs, one positive clone (DNA30942) was identified. When searching for potential alternatively spliced forms of DcR3 that might encode a transmembrane protein, we isolated 50 more clones; the coding regions of these clones were identical in size to that of the initial clone (data not shown).

Fc-fusion proteins (immunoadhesins). The entire DcR3 sequence, or the ectodomain of Fas or TNFR1, was fused to the hinge and Fc region of human IgG1, expressed in insect SF9 cells or in human 293 cells, and purified as described²³.

Fluorescence-activated cell sorting (FACS) analysis. We transfected 293 cells using calcium phosphate or Effectene (Qiagen) with pRK5 vector or pRK5 encoding full-length human FasL⁴ (2 µg), together with pRK5 encoding CrmA (2 µg) to prevent cell death. After 16 h, the cells were incubated with biotinylated DcR3-Fc or TNFR1-Fc and then with phycoerythrin-conjugated streptavidin (GibcoBRL), and were assayed by FACS. The data were analysed by Kolmogorov-Smirnov statistical analysis. There was some detectable staining of vector-transfected cells by DcR3-Fc; as these cells express little FasL (data not shown), it is possible that DcR3 recognized some other factor that is expressed constitutively on 293 cells.

Immunoprecipitation. Human 293 cells were transfected as above, and metabolically labelled with [³⁵S]cysteine and [³⁵S]methionine (0.5 mCi; Amersham). After 16 h of culture in the presence of z-VAD-fmk (10 µM), the medium was immunoprecipitated with DcR3-Fc, Fas-Fc or TNFR1-Fc (5 µg), followed by protein A-Sepharose (Repligen). The precipitates were resolved by SDS-PAGE and visualized on a phosphorimager (Fuji BAS2000). Alternatively, purified, Flag-tagged soluble FasL (1 µg) (Alexis) was incubated with each Fc-fusion protein (1 µg), precipitated with protein A-Sepharose, resolved by SDS-PAGE and visualized by immunoblotting with rabbit anti-FasL antibody (Oncogene Research).

Analysis of complex formation. Flag-tagged soluble FasL (25 µg) was incubated with buffer or with DcR3-Fc (40 µg) for 1.5 h at 24 °C. The reaction was loaded onto a Superdex 200 HR 10/30 column (Pharmacia) and developed with PBS; 0.6-ml fractions were collected. The presence of DcR3-Fc-FasL complex in each fraction was analysed by placing 100 µl aliquots into microtitre wells precoated with anti-human IgG (Boehringer) to capture DcR3-Fc, followed by detection with biotinylated anti-Flag antibody Bio M2 (Kodak) and streptavidin-horseradish peroxidase (Amersham). Calibration of the column indicated an apparent relative molecular mass of the complex of 420K (data not shown), which is consistent with a stoichiometry of two DcR3-Fc homodimers to two soluble FasL homotrimers.

Equilibrium binding analysis. Microtitre wells were coated with anti-human

IgG, blocked with 2% BSA in PBS. DcR3-Fc or Fas-Fc was added, followed by serially diluted Flag-tagged soluble FasL. Bound ligand was detected with anti-Flag antibody as above. In the competition assay, Fas-Fc was immobilized as above; and the wells were blocked with excess IgG1 before addition of Flag-tagged soluble FasL plus DcR3-Fc.

T-cell AICD. CD3⁺ lymphocytes were isolated from peripheral blood of individual donors using anti-CD3 magnetic beads (Miltenyi Biotech), stimulated with phytohaemagglutinin (PHA; 2 µg ml⁻¹) for 24 h, and cultured in the presence of interleukin-2 (100 U ml⁻¹) for 5 days. The cells were plated in wells coated with anti-CD3 antibody (Pharmingen) and analysed for apoptosis 16 h later by FACS analysis of annexin-V-binding of CD4⁺ cells²⁴.

Natural killer cell activity. Natural killer cells were isolated from peripheral blood of individual donors using anti-CD56 magnetic beads (Miltenyi Biotech), and incubated for 16 h with ⁵¹Cr-loaded Jurkat cells at an effector-to-target ratio of 1:1 in the presence of DcR3-Fc, Fas-Fc or human IgG1. Target-cell death was determined by release of ⁵¹Cr in effector-target co-cultures relative to release of ⁵¹Cr by detergent lysis of equal numbers of Jurkat cells.

Gene-amplification analysis. Surgical specimens were provided by J. Kern (lung tumours) and P. Quirke (colon tumours). Genomic DNA was extracted (Qiagen) and the concentration was determined using Hoechst dye 33258 intercalation fluorometry. Amplification was determined by quantitative PCR¹⁸ using a TaqMan instrument (ABI). The method was validated by comparison of PCR and Southern hybridization data for the Myc and HER-2 oncogenes (data not shown). Gene-specific primers and fluorogenic probes were designed on the basis of the sequence of DcR3 or of nearby regions identified on a BAC carrying the human DcR3 gene; alternatively, primers and probes were based on Stanford Human Genome Center marker AFM218xe7 (T160), which is linked to DcR3 (likelihood score = 5.4), SHGC-36268 (T159), the nearest available marker which maps to ~500 kilobases from T160, and five extra markers that span chromosome 20. The DcR3-specific primer sequences were 5'-CTTCTTCGCGCAGCTG-3' and 5'-ATCAGCCGCGCACCAG-3' and the fluorogenic probe sequence was 5'-(FAM-ACACGATGCGTGTCCACGAG AAp-(TAMARA), where FAM is 5'-fluorescein phosphoramidite. Relative gene-copy numbers were derived using the formula 2^(ΔCT), where ΔCT is the difference in amplification cycles required to detect DcR3 in peripheral blood lymphocyte DNA compared to test DNA.

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Correspondence and requests for materials should be addressed to A.A. (e-mail: aa@gene.com). The GenBank accession number for the DcR3 cDNA sequence is AF104419.

Crystal structure of the ATP-binding subunit of an ABC transporter

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ABC transporters (also known as traffic ATPases) form a large family of proteins responsible for the translocation of a variety of compounds across membranes of both prokaryotes and eukaryotes¹. The recently completed *Escherichia coli* genome sequence revealed that the largest family of paralogous *E. coli* proteins is composed of ABC transporters². Many eukaryotic proteins of medical significance belong to this family, such as the cystic fibrosis transmembrane conductance regulator (CFTR), the P-glycoprotein (or multidrug-resistance protein) and the heterodimeric transporter associated with antigen processing (Tap1–Tap2). Here we report the crystal structure at 1.5 Å resolution of HisP, the ATP-binding subunit of the histidine permease, which is an ABC transporter from *Salmonella typhimurium*. We correlate the details of this structure with the biochemical, genetic and biophysical properties of the wild-type and several mutant HisP proteins. The structure provides a basis for understanding properties of ABC transporters and of defective CFTR proteins.

ABC transporters contain four structural domains: two nucleotide-binding domains (NBDs), which are highly conserved throughout the family, and two transmembrane domains¹. In prokaryotes these domains are often separate subunits which are assembled into a membrane-bound complex; in eukaryotes the domains are generally fused into a single polypeptide chain. The periplasmic histidine permease of *S. typhimurium* and *E. coli*^{3–8} is a well-characterized ABC transporter that is a good model for this superfamily. It consists of a membrane-bound complex, HisQMP₂, which comprises integral membrane subunits, HisQ and HisM, and two copies of HisP, the ATP-binding subunit. HisP, which has properties intermediate between those of integral and peripheral membrane proteins⁹, is accessible from both sides of the membrane, presumably by its interaction with HisQ and HisM⁶. The two HisP subunits form a dimer, as shown by their cooperativity in ATP hydrolysis⁵, the requirement for both subunits to be present for activity⁷, and the formation of a HisP dimer upon chemical cross-linking. Soluble HisP also forms a dimer¹. HisP has been purified and characterized in an active soluble form¹ which can be reconstituted into a fully active membrane-bound complex¹.

The overall shape of the crystal structure of the HisP monomer is that of an 'L' with two thick arms (arm I and arm II); the ATP-binding pocket is near the end of arm I (Fig. 1). A six-stranded β -sheet ($\beta 3$ and $\beta 8$ – $\beta 12$) spans both arms of the L, with a domain of α - plus β -type structure ($\beta 1$, $\beta 2$, $\beta 4$ – $\beta 7$, $\alpha 1$ and $\alpha 2$) on one side (within arm I) and a domain of mostly α -helices ($\alpha 3$ – $\alpha 9$) on the

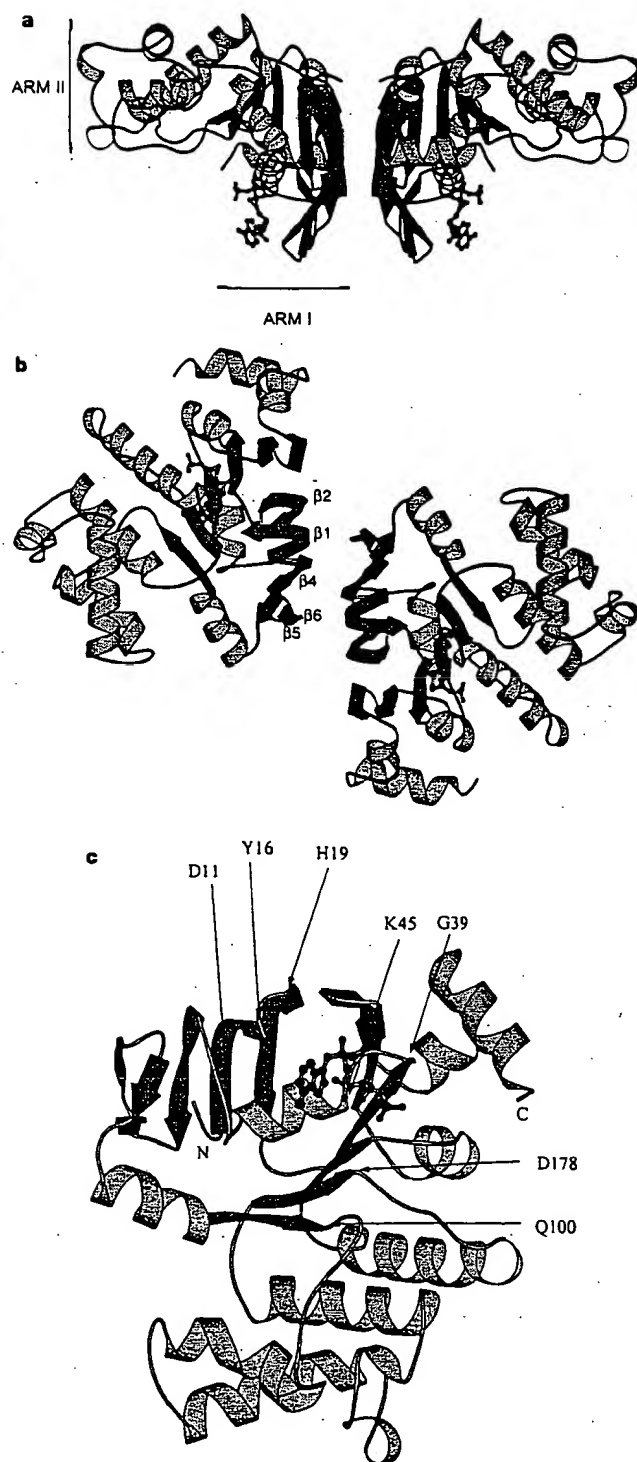


Figure 1 Crystal structure of HisP. **a**, View of the dimer along an axis perpendicular to its two-fold axis. The top and bottom of the dimer are suggested to face towards the periplasmic and cytoplasmic sides, respectively (see text). The thickness of arm II is about 25 Å, comparable to that of membrane. α -Helices are shown in orange and β -sheets in green. **b**, View along the two-fold axis of the HisP dimer, showing the relative displacement of the monomers not apparent in **a**. The β -strands at the dimer interface are labelled. **c**, View of one monomer from the bottom of arm I, as shown in **a**, towards arm II, showing the ATP-binding pocket. **a–c**, The protein and the bound ATP are in 'ribbon' and 'ball-and-stick' representations, respectively. Key residues discussed in the text are indicated in **c**. These figures were prepared with MOLSCRIPT. N, amino terminus; C, C terminus.



NOVEL APPROACH TO QUANTITATIVE POLYMERASE CHAIN REACTION USING REAL-TIME DETECTION: APPLICATION TO THE DETECTION OF GENE AMPLIFICATION IN BREAST CANCER

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Gene amplification is a common event in the progression of human cancers, and amplified oncogenes have been shown to have diagnostic, prognostic and therapeutic relevance. A kinetic quantitative polymerase-chain-reaction (PCR) method, based on fluorescent TaqMan methodology and a new instrument (ABI Prism 7700 Sequence Detection System) capable of measuring fluorescence in real-time, was used to quantify gene amplification in tumor DNA. Reactions are characterized by the point during cycling when PCR amplification is still in the exponential phase, rather than the amount of PCR product accumulated after a fixed number of cycles. None of the reaction components is limited during the exponential phase, meaning that values are highly reproducible in reactions starting with the same copy number. This greatly improves the precision of DNA quantification. Moreover, real-time PCR does not require post-PCR sample handling, thereby preventing potential PCR-product carry-over contamination; it possesses a wide dynamic range of quantification and results in much faster and higher sample throughput. The real-time PCR method, was used to develop and validate a simple and rapid assay for the detection and quantification of the 3 most frequently amplified genes (*myc*, *ccnd1* and *erbB2*) in breast tumors. Extra copies of *myc*, *ccnd1* and *erbB2* were observed in 10, 23 and 15%, respectively, of 108 breast-tumor DNA; the largest observed numbers of gene copies were 4.6, 18.6 and 15.1, respectively. These results correlated well with those of Southern blotting. The use of this new semi-automated technique will make molecular analysis of human cancers simpler and more reliable, and should find broad applications in clinical and research settings. *Int. J. Cancer* 78:661–666, 1998.

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Gene amplification plays an important role in the pathogenesis of various solid tumors, including breast cancer, probably because over-expression of the amplified target genes confers a selective advantage. The first technique used to detect genomic amplification was cytogenetic analysis. Amplification of several chromosome regions, visualized either as extrachromosomal double minutes (dmins) or as integrated homogeneously staining regions (HSRs), are among the main visible cytogenetic abnormalities in breast tumors. Other techniques such as comparative genomic hybridization (CGH) (Kallioniemi *et al.*, 1994) have also been used in broad searches for regions of increased DNA copy numbers in tumor cells, and have revealed some 20 amplified chromosome regions in breast tumors. Positional cloning efforts are underway to identify the critical gene(s) in each amplified region. To date, genes known to be amplified frequently in breast cancers include *myc* (8q24), *ccnd1* (11q13), and *erbB2* (17q12–q21) (for review, see Bièche and Lidereau, 1995).

Amplification of the *myc*, *ccnd1*, and *erbB2* proto-oncogenes should have clinical relevance in breast cancer, since independent studies have shown that these alterations can be used to identify sub-populations with a worse prognosis (Berns *et al.*, 1992; Schuurin *et al.*, 1992; Slamon *et al.*, 1987). Muss *et al.* (1994) suggested that these gene alterations may also be useful for the prediction and assessment of the efficacy of adjuvant chemotherapy and hormone therapy.

However, published results diverge both in terms of the frequency of these alterations and their clinical value. For instance, over 500 studies in 10 years have failed to resolve the controversy

surrounding the link suggested by Slamon *et al.* (1987) between *erbB2* amplification and disease progression. These discrepancies are partly due to the clinical, histological and ethnic heterogeneity of breast cancer, but technical considerations are also probably involved.

Specific genes (DNA) were initially quantified in tumor cells by means of blotting procedures such as Southern and slot blotting. These batch techniques require large amounts of DNA (5–10 µg/reaction) to yield reliable quantitative results. Furthermore, meticulous care is required at all stages of the procedures to generate blots of sufficient quality for reliable dosage analysis. Recently, PCR has proven to be a powerful tool for quantitative DNA analysis, especially with minimal starting quantities of tumor samples (small, early-stage tumors and formalin-fixed, paraffin-embedded tissues).

Quantitative PCR can be performed by evaluating the amount of product either after a given number of cycles (end-point quantitative PCR) or after a varying number of cycles during the exponential phase (kinetic quantitative PCR). In the first case, an internal standard distinct from the target molecule is required to ascertain PCR efficiency. The method is relatively easy but implies generating, quantifying and storing an internal standard for each gene studied. Nevertheless, it is the most frequently applied method to date.

One of the major advantages of the kinetic method is its rapidity in quantifying a new gene, since no internal standard is required (an external standard curve is sufficient). Moreover, the kinetic method has a wide dynamic range (at least 5 orders of magnitude), giving an accurate value for samples differing in their copy number. Unfortunately, the method is cumbersome and has therefore been rarely used. It involves aliquot sampling of each assay mix at regular intervals and quantifying, for each aliquot, the amplification product. Interest in the kinetic method has been stimulated by a novel approach using fluorescent TaqMan methodology and a new instrument (ABI Prism 7700 Sequence Detection System) capable of measuring fluorescence in real time (Gibson *et al.*, 1996; Heid *et al.*, 1996). The TaqMan reaction is based on the 5' nuclease assay first described by Holland *et al.* (1991). The latter uses the 5' nuclease activity of Taq polymerase to cleave a specific fluorogenic oligonucleotide probe during the extension phase of PCR. The approach uses dual-labeled fluorogenic hybridization probes (Lee *et al.*, 1993). One fluorescent dye, co-valently linked to the 5' end of the oligonucleotide, serves as a reporter [FAM (*i.e.*, 6-carboxy-fluorescein)] and its emission spectrum is quenched by a second fluorescent dye, TAMRA (*i.e.*, 6-carboxy-tetramethyl-rhodamine) attached to the 3' end. During the extension phase of the PCR

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cycle, the fluorescent hybridization probe is hydrolyzed by the 5'-3' nucleolytic activity of DNA polymerase. Nuclease degradation of the probe releases the quenching of FAM fluorescence emission, resulting in an increase in peak fluorescence emission. The fluorescence signal is normalized by dividing the emission intensity of the reporter dye (FAM) by the emission intensity of a reference dye (i.e., ROX, 6-carboxy-X-rhodamine) included in TaqMan buffer, to obtain a ratio defined as the R_n (normalized reporter) for a given reaction tube. The use of a sequence detector enables the fluorescence spectra of all 96 wells of the thermal cycler to be measured continuously during PCR amplification.

The real-time PCR method offers several advantages over other current quantitative PCR methods (Celi *et al.*, 1994): (i) the probe-based homogeneous assay provides a real-time method for detecting only specific amplification products, since specific hybridization of both the primers and the probe is necessary to generate a signal; (ii) the C_t (threshold cycle) value used for quantification is measured when PCR amplification is still in the log phase of PCR product accumulation. This is the main reason why C_t is a more reliable measure of the starting copy number than are end-point measurements, in which a slight difference in a limiting component can have a drastic effect on the amount of product; (iii) use of C_t values gives a wider dynamic range (at least 5 orders of magnitude), reducing the need for serial dilution; (iv) The real-time PCR method is run in a closed-tube system and requires no post-PCR sample handling, thus avoiding potential contamination; (v) the system is highly automated, since the instrument continuously measures fluorescence in all 96 wells of the thermal cycler during PCR amplification and the corresponding software processes, and analyzes the fluorescence data; (vi) the assay is rapid, as results are available just one minute after thermal cycling is complete; (vii) the sample throughput of the method is high, since 96 reactions can be analyzed in 2 hr.

Here, we applied this semi-automated procedure to determine the copy numbers of the 3 most frequently amplified genes in breast tumors (*myc*, *ccnd1* and *erbB2*), as well as 2 genes (*alb* and *app*) located in a chromosome region in which no genetic changes have been observed in breast tumors. The results for 108 breast tumors were compared with previous Southern-blot data for the same samples.

MATERIAL AND METHODS

Tumor and blood samples

Samples were obtained from 108 primary breast tumors removed surgically from patients at the Centre René Huguénin; none of the patients had undergone radiotherapy or chemotherapy. Immediately after surgery, the tumor samples were placed in liquid nitrogen until extraction of high-molecular-weight DNA. Patients were included in this study if the tumor sample used for DNA preparation contained more than 60% of tumor cells (histological analysis). A blood sample was also taken from 18 of the same patients.

DNA was extracted from tumor tissue and blood leukocytes according to standard methods.

Real-time PCR

Theoretical basis. Reactions are characterized by the point during cycling when amplification of the PCR product is first detected, rather than by the amount of PCR product accumulated after a fixed number of cycles. The higher the starting copy number of the genomic DNA target, the earlier a significant increase in fluorescence is observed. The parameter C_t (threshold cycle) is defined as the fractional cycle number at which the fluorescence generated by cleavage of the probe passes a fixed threshold above baseline. The target gene copy number in unknown samples is quantified by measuring C_t and by using a standard curve to determine the starting copy number. The precise amount of genomic DNA (based on optical density) and its quality (i.e., lack

of extensive degradation) are both difficult to assess. We therefore also quantified a control gene (*alb*) mapping to chromosome region 4q11-q13, in which no genetic alterations have been found in breast-tumor DNA by means of CGH (Kallioniemi *et al.*, 1994).

Thus, the ratio of the copy number of the target gene to the copy number of the *alb* gene normalizes the amount and quality of genomic DNA. The ratio defining the level of amplification is termed "N", and is determined as follows:

$$N = \frac{\text{copy number of target gene (app, myc, ccnd1, erbB2)}}{\text{copy number of reference gene (alb)}}$$

Primers, probes, reference human genomic DNA and PCR consumables. Primers and probes were chosen with the assistance of the computer programs Oligo 4.0 (National Biosciences, Plymouth, MN), EuGene (Daniben Systems, Cincinnati, OH) and Primer Express (Perkin-Elmer Applied Biosystems, Foster City, CA).

Primers were purchased from DNAgency (Malvern, PA) and probes from Perkin-Elmer Applied Biosystems.

Nucleotide sequences for the oligonucleotide hybridization probes and primers are available on request.

The TaqMan PCR Core reagent kit, MicroAmp optical tubes, and MicroAmp caps were from Perkin-Elmer Applied Biosystems.

Standard-curve construction. The kinetic method requires a standard curve. The latter was constructed with serial dilutions of specific PCR products, according to Piatak *et al.* (1993). In practice, each specific PCR product was obtained by amplifying 20 ng of a standard human genomic DNA (Boehringer, Mannheim, Germany) with the same primer pairs as those used later for real-time quantitative PCR. The 5 PCR products were purified using MicroSpin S-400 HR columns (Pharmacia, Uppsala, Sweden) electrophoresed through an acrylamide gel and stained with ethidium bromide to check their quality. The PCR products were then quantified spectrophotometrically and pooled, and serially diluted 10-fold in mouse genomic DNA (Clontech, Palo Alto, CA) at a constant concentration of 2 ng/ μ l. The standard curve used for real-time quantitative PCR was based on serial dilutions of the pool of PCR products ranging from 10^{-7} (10^5 copies of each gene) to 10^{-10} (10^2 copies). This series of diluted PCR products was aliquoted and stored at -80°C until use.

The standard curve was validated by analyzing 2 known quantities of calibrator human genomic DNA (20 ng and 50 ng).

PCR amplification. Amplification mixes (50 μ l) contained the sample DNA (around 20 ng, around 6600 copies of disomic genes), $10\times$ TaqMan buffer (5 μ l), 200 μ M dATP, dCTP, dGTP, and 400 μ M dUTP, 5 mM MgCl_2 , 1.25 units of AmpliTaq Gold, 0.5 units of AmpErase uracil N-glycosylase (UNG), 200 nM each primer and 100 nM probe. The thermal cycling conditions comprised 2 min at 50°C and 10 min at 95°C . Thermal cycling consisted of 40 cycles at 95°C for 15 s and 65°C for 1 min. Each assay included: a standard curve (from 10^5 to 10^2 copies) in duplicate, a no-template control, 20 ng and 50 ng of calibrator human genomic DNA (Boehringer) in triplicate, and about 20 ng of unknown genomic DNA in triplicate (26 samples can thus be analyzed on a 96-well microplate). All samples with a coefficient of variation (CV) higher than 10% were retested.

All reactions were performed in the ABI Prism 7700 Sequence Detection System (Perkin-Elmer Applied Biosystems), which detects the signal from the fluorogenic probe during PCR.

Equipment for real-time detection. The 7700 system has a built-in thermal cycler and a laser directed via fiber optical cables to each of the 96 sample wells. A charge-coupled-device (CDD) camera collects the emission from each sample and the data are analyzed automatically. The software accompanying the 7700 system calculates C_t and determines the starting copy number in the samples.

Determination of gene amplification. Gene amplification was calculated as described above. Only samples with an N value higher than 2 were considered to be amplified.

RESULTS

To validate the method, real-time PCR was performed on genomic DNA extracted from 108 primary breast tumors, and 18 normal leukocyte DNA samples from some of the same patients. The target genes were the *myc*, *ccnd1* and *erbB2* proto-oncogenes, and the β -amyloid precursor protein gene (*app*), which maps to a chromosome region (21q21.2) in which no genetic alterations have been found in breast tumors (Kallioniemi *et al.*, 1994). The reference disomic gene was the albumin gene (*alb*, chromosome 4q11-q13).

Validation of the standard curve and dynamic range of real-time PCR

The standard curve was constructed from PCR products serially diluted in genomic mouse DNA at a constant concentration of 2 ng/ μ l. It should be noted that the 5 primer pairs chosen to analyze the 5 target genes do not amplify genomic mouse DNA (data not shown). Figure 1 shows the real-time PCR standard curve for the *alb* gene. The dynamic range was wide (at least 4 orders of magnitude), with samples containing as few as 10^2 copies or as many as 10^5 copies.

Copy-number ratio of the 2 reference genes (*app* and *alb*)

The *app* to *alb* copy-number ratio was determined in 18 normal leukocyte DNA samples and all 108 primary breast-tumor DNA

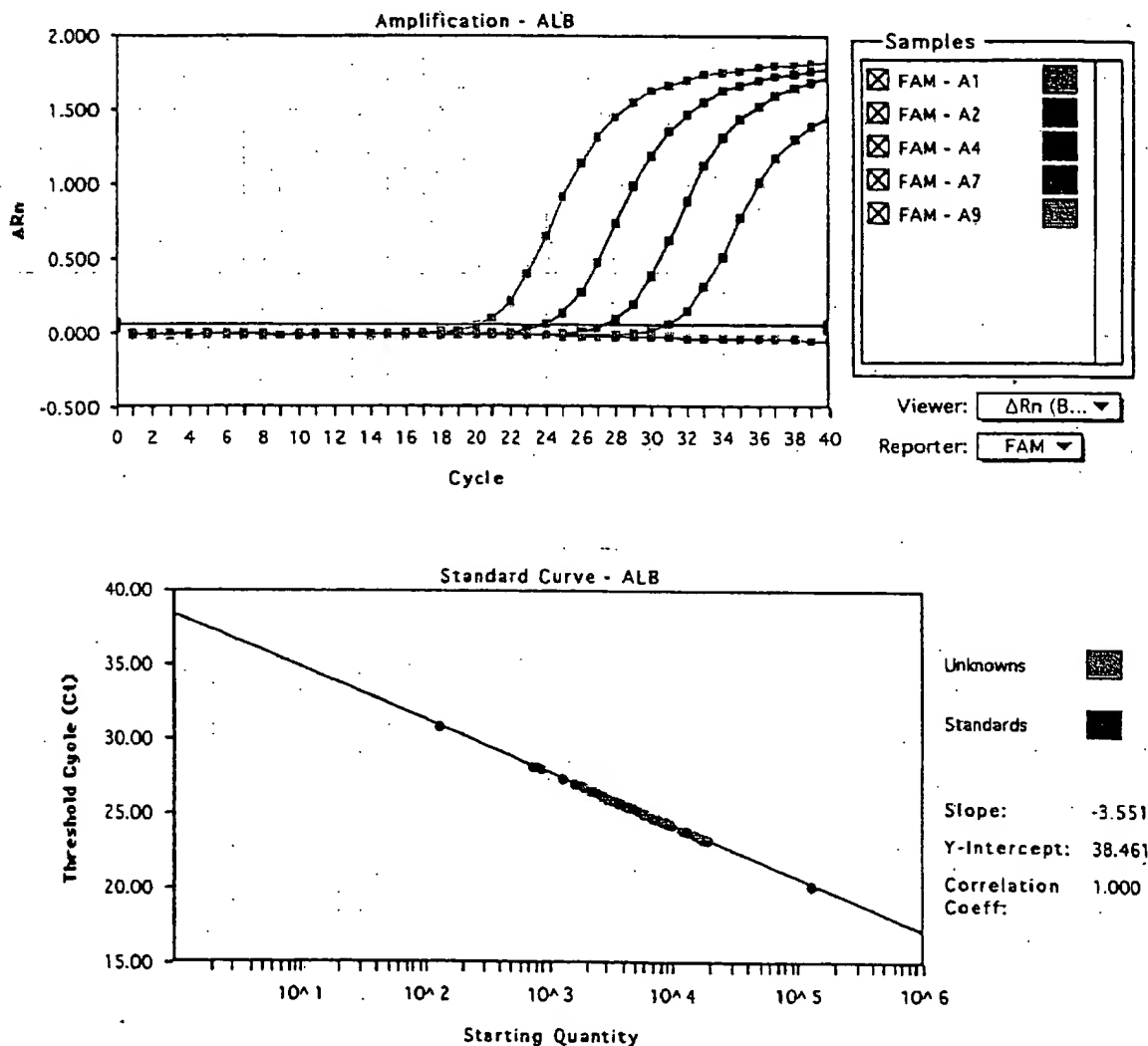


FIGURE 1 – Albumin (*alb*) gene dosage by real-time PCR. Top: Amplification plots for reactions with starting *alb* gene copy number ranging from 10^5 (A9), 10^4 (A7), 10^3 (A4) to 10^2 (A2) and a no-template control (A1). Cycle number is plotted vs. change in normalized reporter signal (ΔRn). For each reaction tube, the fluorescence signal of the reporter dye (FAM) is divided by the fluorescence signal of the passive reference dye (ROX), to obtain a ratio defined as the normalized reporter signal (Rn). ΔRn represents the normalized reporter signal (Rn) minus the baseline signal established in the first 15 PCR cycles. ΔRn increases during PCR as *alb* PCR product copy number increases until the reaction reaches a plateau. C_t (threshold cycle) represents the fractional cycle number at which a significant increase in Rn above a baseline signal (horizontal black line) can first be detected. Two replicate plots were performed for each standard sample, but the data for only one are shown here. Bottom: Standard curve plotting log starting copy number vs. C_t (threshold cycle). The black dots represent the data for standard samples plotted in duplicate and the red dots the data for unknown genomic DNA samples plotted in triplicate. The standard curve shows 4 orders of linear dynamic range.

samples. We selected these 2 genes because they are located in 2 chromosome regions (*app*, 21q21.2; *alb*, 4q11-q13) in which no obvious genetic changes (including gains or losses) have been observed in breast cancers (Kallioniemi *et al.*, 1994). The ratio for the 18 normal leukocyte DNA samples fell between 0.7 and 1.3 (mean 1.02 ± 0.21), and was similar for the 108 primary breast-tumor DNA samples (0.6 to 1.6 , mean 1.06 ± 0.25), confirming that *alb* and *app* are appropriate reference disomic genes for breast-tumor DNA. The low range of the ratios also confirmed that the nucleotide sequences chosen for the primers and probes were not polymorphic, as mismatches of their primers or probes with the subject's DNA would have resulted in differential amplification.

myc, *ccnd1* and *erbB2* gene dose in normal leukocyte DNA

To determine the cut-off point for gene amplification in breast-cancer tissue, 18 normal leukocyte DNA samples were tested for the gene dose (N), calculated as described in "Material and Methods". The N value of these samples ranged from 0.5 to 1.3 (mean 0.84 ± 0.22) for *myc*, 0.7 to 1.6 (mean 1.06 ± 0.23) for *ccnd1* and 0.6 to 1.3 (mean 0.91 ± 0.19) for *erbB2*. Since N values for *myc*, *ccnd1* and *erbB2* in normal leukocyte DNA consistently fell between 0.5 and 1.6, values of 2 or more were considered to represent gene amplification in tumor DNA.

myc, *ccnd1* and *erbB2* gene dose in breast-tumor DNA

myc, *ccnd1* and *erbB2* gene copy numbers in the 108 primary breast tumors are reported in Table I. Extra copies of *ccnd1* were more frequent (23%, 25/108) than extra copies of *erbB2* (15%, 16/108) and *myc* (10%, 11/108), and ranged from 2 to 18.6 for *ccnd1*, 2 to 15.1 for *erbB2*, and only 2 to 4.6 for the *myc* gene. Figure 2 and Table II represent tumors in which the *ccnd1* gene was amplified 16-fold (T145), 6-fold (T133) and non-amplified (T118). The 3 genes were never found to be co-amplified in the same tumor. *erbB2* and *ccnd1* were co-amplified in only 3 cases, *myc* and *ccnd1* in 2 cases and *myc* and *erbB2* in 1 case. This favors the hypothesis that gene amplifications are independent events in breast cancer. Interestingly, 5 tumors showed a decrease of at least 50% in the *erbB2* copy number ($N < 0.5$), suggesting that they bore deletions of the 17q21 region (the site of *erbB2*). No such decrease in copy number was observed with the other 2 proto-oncogenes.

Comparison of gene dose determined by real-time quantitative PCR and Southern-blot analysis

Southern-blot analysis of *myc*, *ccnd1* and *erbB2* amplifications had previously been done on the same 108 primary breast tumors. A perfect correlation between the results of real-time PCR and Southern blot was obtained for tumors with high copy numbers ($N \geq 5$). However, there were cases (1 *myc*, 6 *ccnd1* and 4 *erbB2*) in which real-time PCR showed gene amplification whereas Southern-blot did not, but these were mainly cases with low extra copy numbers (N from 2 to 2.9).

DISCUSSION

The clinical applications of gene amplification assays are currently limited, but would certainly increase if a simple, standardized and rapid method were perfected. Gene amplification status has been studied mainly by means of Southern blotting, but this method is not sensitive enough to detect low-level gene amplification nor accurate enough to quantify the full range of amplification values. Southern blotting is also time-consuming, uses radioactive

reagents and requires relatively large amounts of high-quality genomic DNA, which means it cannot be used routinely in many laboratories. An amplification step is therefore required to determine the copy number of a given target gene from minimal quantities of tumor DNA (small early-stage tumors, cytopuncture specimens or formalin-fixed, paraffin-embedded tissues).

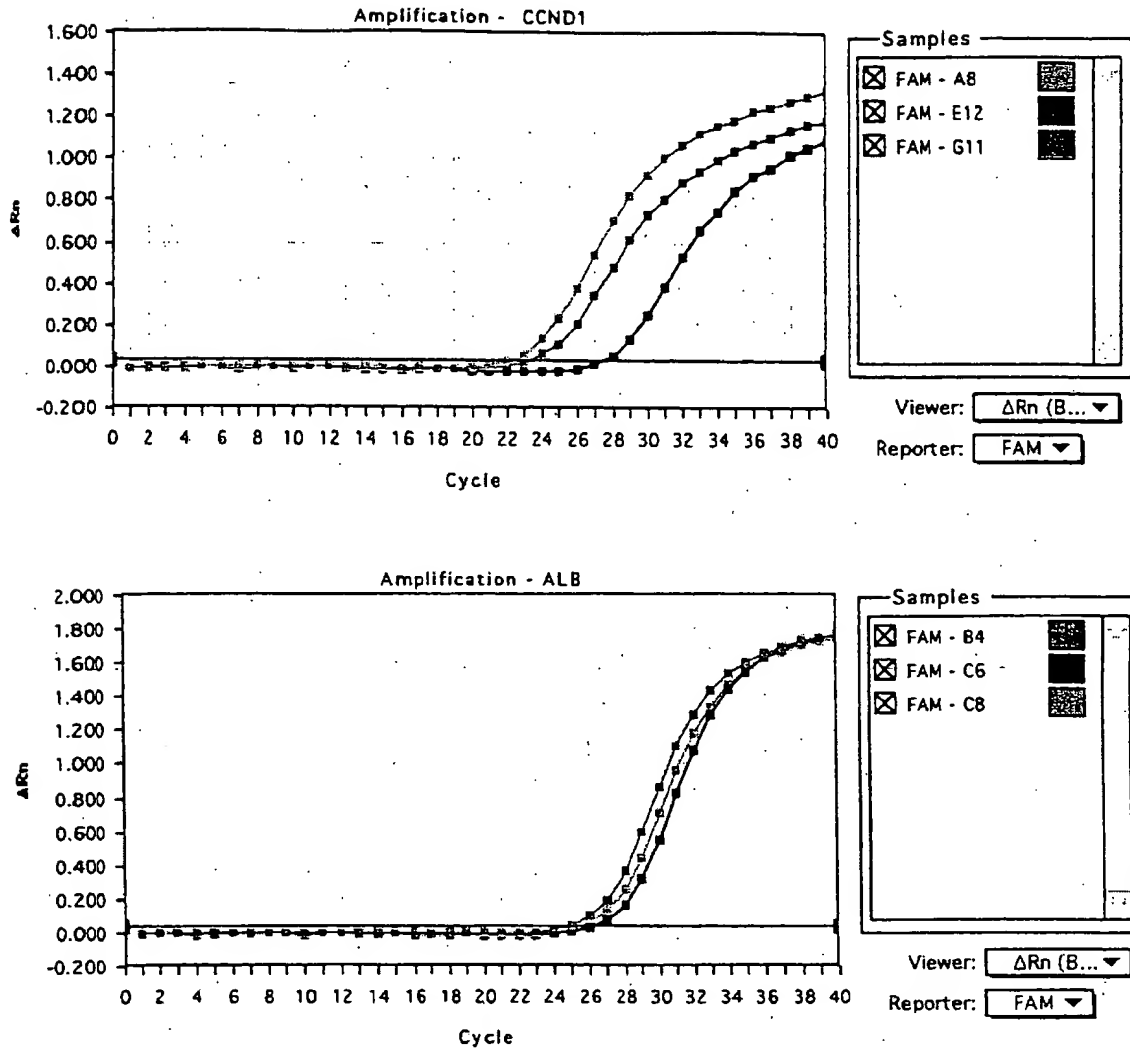
In this study, we validated a PCR method developed for the quantification of gene over-representation in tumors. The method, based on real-time analysis of PCR amplification, has several advantages over other PCR-based quantitative assays such as competitive quantitative PCR (Celi *et al.*, 1994). First, the real-time PCR method is performed in a closed-tube system, avoiding the risk of contamination by amplified products. Re-amplification of carryover PCR products in subsequent experiments can also be prevented by using the enzyme uracil N-glycosylase (UNG) (Longo *et al.*, 1990). The second advantage is the simplicity and rapidity of sample analysis, since no post-PCR manipulations are required. Our results show that the automated method is reliable. We found it possible to determine, in triplicate, the number of copies of a target gene in more than 100 tumors per day. Third, the system has a linear dynamic range of at least 4 orders of magnitude, meaning that samples do not have to contain equal starting amounts of DNA. This technique should therefore be suitable for analyzing formalin-fixed, paraffin-embedded tissues. Fourth, and above all, real-time PCR makes DNA quantification much more precise and reproducible, since it is based on C_t values rather than end-point measurement of the amount of accumulated PCR product. Indeed, the ABI Prism 7700 Sequence Detection System enables C_t to be calculated when PCR amplification is still in the exponential phase and when none of the reaction components is rate-limiting. The within-run CV of the C_t value for calibrator human DNA (5 replicates) was always below 5%, and the between-assay precision in 5 different runs was always below 10% (data not shown). In addition, the use of a standard curve is not absolutely necessary, since the copy number can be determined simply by comparing the C_t ratio of the target gene with that of reference genes. The results obtained by the 2 methods (with and without a standard curve) are similar in our experiments (data not shown). Moreover, unlike competitive quantitative PCR, real-time PCR does not require an internal control (the design and storage of internal controls and the validation of their amplification efficiency is laborious).

The only potential disadvantage of real-time PCR, like all other PCR-based methods and solid-matrix blotting techniques (Southern blots and dot blots) is that it cannot avoid dilution artifacts inherent in the extraction of DNA from tumor cells contained in heterogeneous tissue specimens. Only FISH and immunohistochemistry can measure alterations on a cell-by-cell basis (Pauletti *et al.*, 1996; Slamon *et al.*, 1989). However, FISH requires expensive equipment and trained personnel and is also time-consuming. Moreover, FISH does not assess gene expression and therefore cannot detect cases in which the gene product is over-expressed in the absence of gene amplification, which will be possible in the future by real-time quantitative RT-PCR. Immunohistochemistry is subject to considerable variations in the hands of different teams, owing to alterations of target proteins during the procedure, the different primary antibodies and fixation methods used and the criteria used to define positive staining.

The results of this study are in agreement with those reported in the literature. (i) Chromosome regions 4q11-q13 and 21q21.2 (which bear *alb* and *app*, respectively) showed no genetic alterations in the breast-cancer samples studied here, in keeping with the results of CGH (Kallioniemi *et al.*, 1994). (ii) We found that amplifications of these 3 oncogenes were independent events, as reported by other teams (Berns *et al.*, 1992; Borg *et al.*, 1992). (iii) The frequency and degree of *myc* amplification in our breast tumor DNA series were lower than those of *ccnd1* and *erbB2* amplification, confirming the findings of Borg *et al.* (1992) and Courjal *et al.* (1997). (iv) The maxima of *ccnd1* and *erbB2* over-representation were 18-fold and 15-fold, also in keeping with earlier results (about

TABLE I - DISTRIBUTION OF AMPLIFICATION LEVEL (N) FOR *myc*, *ccnd1* AND *erbB2* GENES IN 108 HUMAN BREAST TUMORS

Gene	Amplification level (N)			
	<0.5	0.5-1.9	2-4.9	≥ 5
<i>myc</i>	0	97 (89.8%)	11 (10.2%)	0
<i>ccnd1</i>	0	83 (76.9%)	17 (15.7%)	8 (7.4%)
<i>erbB2</i>	5 (4.6%)	87 (80.6%)	8 (7.4%)	8 (7.4%)



Tumor	CCND1		ALB	
	C_t	Copy number	C_t	Copy number
T118	27.3	4605	26.5	4365
T133	23.2	61659	25.2	10092
T145	22.1	125892	25.6	7762

FIGURE 2 - *ccnd1* and *alb* gene dosage by real-time PCR in 3 breast tumor samples: T118 (E12, C6, black squares), T133 (G11, B4, red squares) and T145 (A8, C8, blue squares). Given the C_t of each sample, the initial copy number is inferred from the standard curve obtained during the same experiment. Triplicate plots were performed for each tumor sample, but the data for only one are shown here. The results are shown in Table II.

30-fold maximum) (Berns *et al.*, 1992; Borg *et al.*, 1992; Courjal *et al.*, 1997). (v) The *erbB2* copy numbers obtained with real-time PCR were in good agreement with data obtained with other quantitative PCR-based assays in terms of the frequency and degree of amplification (An *et al.*, 1995; Deng *et al.*, 1996; Valeron

et al., 1996). Our results also correlate well with those recently published by Gelmini *et al.* (1997), who used the TaqMan system to measure *erbB2* amplification in a small series of breast tumors ($n = 25$), but with an instrument (LS-50B luminescence spectrometer, Perkin-Elmer Applied Biosystems) which only allows end-

TABLE II - EXAMPLES OF *ccnd1* GENE DOSAGE RESULTS FROM 3 BREAST TUMORS¹

Tumor	<i>ccnd1</i>			<i>alb</i>			<i>Nccnd1/alb</i>
	Copy number	Mean	SD	Copy number	Mean	SD	
T118	4525	4603	77	4223	4325	89	1.06
	4605			4365			
	4678			4387			
T133	59821	61100	1111	9787	10137	375	6.03
	61659			10092			
	61821			10533			
T145	128563	125392	3448	7321	7672	316	16.34
	125892			7762			
	121722			7933			

¹For each sample, 3 replicate experiments were performed and the mean and the standard deviation (SD) was determined. The level of *ccnd1* gene amplification (*Nccnd1/alb*) is determined by dividing the average *ccnd1* copy number value by the average *alb* copy number value.

point measurement of fluorescence intensity. Here we report *myc* and *ccnd1* gene dosage in breast cancer by means of quantitative PCR. (vi) We found a high degree of concordance between real-time quantitative PCR and Southern blot analysis in terms of gene amplification, especially for samples with high copy numbers (≥ 5 -fold). The slightly higher frequency of gene amplification (especially *ccnd1* and *erbB2*) observed by means of real-time quantitative PCR as compared with Southern-blot analysis may be explained by the higher sensitivity of the former method. However, we cannot rule out the possibility that some tumors with a few extra

gene copies observed in real-time PCR had additional copies of an arm or a whole chromosome (trisomy, tetrasomy or polysomy) rather than true gene amplification. These 2 types of genetic alteration (polysomy and gene amplification) could be easily distinguished in the future by using an additional probe located on the same chromosome arm, but some distance from the target gene. It is noteworthy that high gene copy numbers have the greatest prognostic significance in breast carcinoma (Borg *et al.*, 1992; Slamon *et al.*, 1987).

Finally, this technique can be applied to the detection of gene deletion as well as gene amplification. Indeed, we found a decreased copy number of *erbB2* (but not of the other 2 proto-oncogenes) in several tumors; *erbB2* is located in a chromosome region (17q21) reported to contain both deletions and amplifications in breast cancer (Bièche and Lidereau, 1995).

In conclusion, gene amplification in various cancers can be used as a marker of pre-neoplasia, also for early diagnosis of cancer, staging, prognostication and choice of treatment. Southern blotting is not sufficiently sensitive, and FISH is lengthy and complex. Real-time quantitative PCR overcomes both these limitations, and is a sensitive and accurate method of analyzing large numbers of samples in a short time. It should find a place in routine clinical gene dosage.

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DECLARATION OF PAUL POLAKIS, Ph.D.

I, Paul Polakis, Ph.D., declare and say as follows:

1. I was awarded a Ph.D. by the Department of Biochemistry of the Michigan State University in 1984. My scientific Curriculum Vitae is attached to and forms part of this Declaration (Exhibit A).
2. I am currently employed by Genentech, Inc. where my job title is Staff Scientist. Since joining Genentech in 1999, one of my primary responsibilities has been leading Genentech's Tumor Antigen Project, which is a large research project with a primary focus on identifying tumor cell markers that find use as targets for both the diagnosis and treatment of cancer in humans.
3. As part of the Tumor Antigen Project, my laboratory has been analyzing differential expression of various genes in tumor cells relative to normal cells. The purpose of this research is to identify proteins that are abundantly expressed on certain tumor cells and that are either (i) not expressed, or (ii) expressed at lower levels, on corresponding normal cells. We call such differentially expressed proteins "tumor antigen proteins". When such a tumor antigen protein is identified, one can produce an antibody that recognizes and binds to that protein. Such an antibody finds use in the diagnosis of human cancer and may ultimately serve as an effective therapeutic in the treatment of human cancer.
4. In the course of the research conducted by Genentech's Tumor Antigen Project, we have employed a variety of scientific techniques for detecting and studying differential gene expression in human tumor cells relative to normal cells, at genomic DNA, mRNA and protein levels. An important example of one such technique is the well known and widely used technique of microarray analysis which has proven to be extremely useful for the identification of mRNA molecules that are differentially expressed in one tissue or cell type relative to another. In the course of our research using microarray analysis, we have identified approximately 200 gene transcripts that are present in human tumor cells at significantly higher levels than in corresponding normal human cells. To date, we have generated antibodies that bind to about 30 of the tumor antigen proteins expressed from these differentially expressed gene transcripts and have used these antibodies to quantitatively determine the level of production of these tumor antigen proteins in both human cancer cells and corresponding normal cells. We have then compared the levels of mRNA and protein in both the tumor and normal cells analyzed.
5. From the mRNA and protein expression analyses described in paragraph 4 above, we have observed that there is a strong correlation between changes in the level of mRNA present in any particular cell type and the level of protein

expressed from that mRNA in that cell type. In approximately 80% of our observations we have found that increases in the level of a particular mRNA correlates with changes in the level of protein expressed from that mRNA when human tumor cells are compared with their corresponding normal cells.

6. Based upon my own experience accumulated in more than 20 years of research, including the data discussed in paragraphs 4 and 5 above and my knowledge of the relevant scientific literature, it is my considered scientific opinion that for human genes, an increased level of mRNA in a tumor cell relative to a normal cell typically correlates to a similar increase in abundance of the encoded protein in the tumor cell relative to the normal cell. In fact, it remains a central dogma in molecular biology that increased mRNA levels are predictive of corresponding increased levels of the encoded protein. While there have been published reports of genes for which such a correlation does not exist, it is my opinion that such reports are exceptions to the commonly understood general rule that increased mRNA levels are predictive of corresponding increased levels of the encoded protein.

7. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: 5/07/04

By: Paul Polakis

Paul Polakis, Ph.D.



①9 BUNDESREPUBLIK
DEUTSCHLAND



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PATENT- UND
MARKENAMT

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Die folgenden Angaben sind den vom Anmelder eingereichten Unterlagen entnommen

- ⑤④ Menschliche Nukleinsäuresequenzen aus Blase-Tumor
⑤⑦ Es werden menschliche Nukleinsäuresequenzen -
mRNA, cDNA, genomische Sequenzen - aus Blasen-
tumorgewebe, die für Genprodukte oder Teile davon kodie-
ren, und deren Verwendung beschrieben.
Es werden weiterhin die über die Sequenzen erhältlichen
Polypeptide und deren Verwendung beschrieben.

DE 198 18 619 A 1

Die Erfindung betrifft menschliche Nukleinsäuresequenzen aus Blasen Tumorgewebe, die für Genprodukte oder Teile davon kodieren, deren funktionale Gene, die mindestens ein biologisch aktives Polypeptid kodieren und deren Verwendung.

Die Erfindung betrifft weiterhin die über die Sequenzen erhältlichen Polypeptide und deren Verwendung.

Eine der Hauptkrebstodesursachen ist der Blasen tumor, für dessen Bekämpfung neue Therapien notwendig sind. Bisher verwendete Therapien, wie z. B. Chemotherapie, Hormontherapie oder chirurgische Entfernung des Tumorgewebes, führen häufig nicht zu einer vollständigen Heilung.

Das Phänomen Krebs geht häufig einher mit der Über- oder Unterexpression gewisser Gene in den entarteten Zellen, wobei noch unklar ist, ob diese veränderten Expressionsraten Ursache oder Folge der malignen Transformation sind. Die Identifikation solcher Gene wäre ein wesentlicher Schritt für die Entwicklung neuer Therapien gegen Krebs. Der spontanen Entstehung von Krebs geht häufig eine Vielzahl von Mutationen voraus. Diese können verschiedenste Auswirkungen auf das Expressionsmuster in dem betroffenen Gewebe haben, wie z. B. Unter- oder Überexpression, aber auch Expression verkürzter Gene. Mehrere solcher Veränderungen durch solche Mutationskaskaden können schließlich zu bösartigen Entartungen führen. Die Komplexität solcher Zusammenhänge erschwert die experimentelle Herangehensweise sehr.

Für die Suche nach Kandidatengen, d. h. Genen, die im Vergleich zum Tumorgewebe im normalen Gewebe stärker exprimiert werden, wird eine Datenbank verwendet, die aus sogenannten ESTs besteht. ESTs (Expressed Sequence Tags) sind Sequenzen von cDNAs, d. h. revers transkribierten mRNAs, den Molekülen also, die die Expression von Genen widerspiegeln. Die EST-Sequenzen werden für normale und entartete Gewebe ermittelt. Solche Datenbanken werden von verschiedenen Betreibern z. T. kommerziell angeboten. Die ESTs der LifeSeq-Datenbank, die hier verwendet wird, sind in der Regel zwischen 150 und 350 Nukleotide lang. Sie repräsentieren ein für ein bestimmtes Gen unverkennbares Muster, obwohl dieses Gen normalerweise sehr viel länger ist (> 2000 Nukleotide). Durch Vergleich der Expressionsmuster von normalen und Tumorgewebe können ESTs identifiziert werden, die für die Tumorentstehung und -proliferation wichtig sind. Es besteht jedoch folgendes Problem: Da durch unterschiedliche Konstruktionen der cDNA-Bibliotheken die gefundenen EST-Sequenzen zu unterschiedlichen Regionen eines unbekannten Gens gehören können, ergäbe sich in einem solchen Fall ein völlig falsches Verhältnis des Vorkommens dieser ESTs in dem jeweiligen Gewebe. Dieses würde erst bemerkt werden, wenn das vollständige Gen bekannt ist und somit die ESTs dem gleichen Gen zugeordnet werden können.

Es wurde nun gefunden, daß diese Fehlermöglichkeit verringert werden kann, wenn zuvor sämtliche ESTs aus dem jeweiligen Gewebstyp assembliert werden, bevor die Expressionsmuster miteinander verglichen werden. Es wurden also überlappende ESTs ein und desselben Gens zu längeren Sequenzen zusammengefaßt (s. Fig. 1, Fig. 2a und Fig. 3). Durch diese Verlängerung und damit Abdeckung eines wesentlich größeren Genbereichs in jeder der jeweiligen Banken sollte der oben beschriebene Fehler weitgehendst vermieden werden. Da es hierzu keine bestehenden Softwareprodukte gab, wurden Programme für das Assemblieren von genomischen Abschnitten verwendet, die abgewandelt eingesetzt und durch eigene Programme ergänzt wurden. Ein Flowchart der Assemblierungsprozedur ist in Fig. 2b1-2b4 dargestellt.

Es konnten nun die Nukleinsäure-Sequenzen Seq. ID No. 1 50 gefunden werden, die als Kandidatengene beim Blasen tumor eine Rolle spielen.

Von besonderem Interesse sind die Nukleinsäure-Sequenzen Seq. ID Nos. 2-5, 7-13, 16, 18, 20, 23, 26-27, 31-32, 36, 45.

Die Erfindung betrifft somit Nukleinsäure-Sequenzen, die ein Genprodukt oder ein Teil davon kodieren, umfassend

- a) eine Nukleinsäure-Sequenz, ausgewählt aus der Gruppe der Nukleinsäure-Sequenzen Seq. ID Nos. 2-5, 7-13, 16, 18, 20, 23, 26-27, 31-32, 36, 45.
- b) eine allelische Variation der unter a) genannten Nukleinsäure-Sequenzen oder
- c) eine Nukleinsäure-Sequenz, die komplementär zu den unter a) oder b) genannten Nukleinsäure-Sequenzen ist.

Die Erfindung betrifft weiterhin eine Nukleinsäure-Sequenz gemäß einer der Sequenzen Seq. ID Nos. 2-5, 7-13, 16, 18, 20, 23, 26-27, 31-32, 36, 45 oder eine komplementäre oder allelische Variante davon und die Nukleinsäure-Sequenzen davon, die eine 90%ige bis 95%ige Homologie zu einer humanen Nukleinsäure-Sequenz aufweisen.

Die Erfindung betrifft auch die Nukleinsäure-Sequenzen Seq. ID No. 1 bis Seq. ID No. 50, die im Blasen tumor Gewebe erhöht exprimiert sind.

Die Erfindung betrifft ferner Nukleinsäure-Sequenzen, umfassend einen Teil der oben genannten Nukleinsäure-Sequenzen, in solch einer ausreichenden Größe, daß sie mit den Sequenzen Seq. ID Nos. 1-50 hybridisieren.

Die erfindungsgemäßen Nukleinsäure-Sequenzen weisen im allgemeinen eine Länge von mindestens 50 bis 4500 bp, vorzugsweise eine Länge von mindestens 150 bis 4000 bp, insbesondere eine Länge von 450 bis 3500 bp auf.

Mit den erfindungsgemäßen Teilsequenzen Seq. ID Nos. 1-50 können gemäß gängiger Verfahrenspraxis auch Expressionskassetten konstruiert werden, wobei auf der Kassette mindestens eine der erfindungsgemäßen Nukleinsäure-Sequenzen zusammen mit mindestens einem dem Fachmann allgemein bekannten Kontroll- oder regulatorischen Sequenz, wie z. B. einem geeigneten Promotor, kombiniert wird. Die erfindungsgemäßen Sequenzen können in sense oder antisense Orientierung eingefügt sein.

In der Literatur sind eine große Anzahl von Expressionskassetten bzw. Vektoren und Promotoren bekannt, die verwendet werden können.

Unter Expressionskassetten bzw. Vektoren sind zu verstehen: 1. bakterielle, wie z. B., phagescript, pBs, ϕ X174, pBluescript SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene), pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia), 2. eukaryontische, wie z. B. pWLneo, pSV2cat, pOG44, pXT1, pSG (Stratagene), pSVK3, pBPV, pMSG,

pSVL (Pharmacia).

Unter Kontroll- oder regulatorischer Sequenz sind geeignete Promotoren zu verstehen. Hierbei sind zwei bevorzugte Vektoren der pKK232-8 und der PCM7 Vektor. Im einzelnen sind folgende Promotoren gemeint: lacI, lacZ, T3, T7, gpt, lambda P_R, trc, CMV, HSV Thymidin-Kinase, SV40, LTRs aus Retrovirus und Maus Metallothioncin-I.

Die auf der Expressionskassette befindlichen DNA-Sequenzen können ein Fusionsprotein kodieren, das ein bekanntes Protein und ein biologisch aktives Polypeptid-Fragment umfaßt. 5

Die Expressionskassetten sind ebenfalls Gegenstand der vorliegenden Erfindung.

Die erfindungsgemäßen Nukleinsäure-Fragmente können zur Herstellung von Vollängen-Genen verwendet werden. Die erhältlichen Gene sind ebenfalls Gegenstand der vorliegenden Erfindung.

Die Erfindung betrifft auch die Verwendung der erfindungsgemäßen Nukleinsäure-Sequenzen, sowie die aus der Verwendung erhältlichen Gen-Fragmente. 10

Die erfindungsgemäßen Nukleinsäure-Sequenzen können mit geeigneten Vektoren in Wirtszellen gebracht werden, in denen als heterologer Teil die auf den Nukleinsäure-Fragmenten enthaltene genetischen Information befindet, die exprimiert wird.

Die die Nukleinsäure-Fragmente enthaltenden Wirtszellen sind ebenfalls Gegenstand der vorliegenden Erfindung. 15

Geeignete Wirtszellen sind z. B. prokaryontische Zellsysteme wie E. coli oder eukaryontische Zellsysteme wie tierische oder humane Zellen oder Hefen.

Die erfindungsgemäßen Nukleinsäure-Sequenzen können in sense oder antisense Form verwendet werden.

Die Herstellung der Polypeptide oder deren Fragment erfolgt durch Kultivierung der Wirtszellen gemäß gängiger Kultivierungsmethoden und anschließender Isolierung und Aufreinigung der Peptide bzw. Fragmente, ebenfalls mittels gängiger Verfahren. Die Erfindung betrifft ferner Nukleinsäure-Sequenzen, die mindestens eine Teilsequenz eines biologisch aktiven Polypeptids kodieren. 20

Ferner betrifft die vorliegende Erfindung Polypeptid-Teilsequenzen, sogenannte ORF (open-reading-frame)-Peptide, gemäß den Sequenzprotokollen ORF ID Nos. 51-106.

Die Erfindung betrifft ferner die Polypeptid-Sequenzen, die mindestens eine 80%ige Homologie, insbesondere eine 90%ige Homologie zu den erfindungsgemäßen Polypeptid-Teilsequenzen der ORF. ID Nos. 51-106 aufweisen. 25

Die Erfindung betrifft auch Antikörper, die gegen ein Polypeptid oder Fragment davon gerichtet sind, welche von den erfindungsgemäßen Nukleinsäuren der Sequenzen Seq. ID No. 1 bis Seq. ID 50 kodiert werden.

Unter Antikörper sind insbesondere monoklonale und Phage-Display-Antikörper zu verstehen.

Die erfindungsgemäßen Polypeptide der Sequenzen ORF ID Nos. 51-106 können auch als Tool zum Auffinden von Wirkstoffen gegen den Blasen-tumor verwendet werden, was ebenfalls Gegenstand der vorliegenden Erfindung ist. 30

Ebenfalls Gegenstand der vorliegenden Erfindung ist die Verwendung der Nukleinsäure-Sequenzen gemäß den Sequenzen Seq. ID No. 1-50 zur Expression von Polypeptiden, die als Tools zum Auffinden von Wirkstoffen gegen den Blasen-tumor verwendet werden können.

Die Erfindung betrifft auch die Verwendung der gefundenen Polypeptid-Teilsequenzen ORF. ID No. 51-106 als Arzneimittel in der Gentherapie zur Behandlung gegen den Blasen-tumor, bzw. zur Herstellung eines Arzneimittels zur Behandlung gegen den Blasen-tumor. 35

Die Erfindung betrifft auch Arzneimittel, die mindestens eine Polypeptid-Teilsequenz ORF. ID No. 51-106 enthalten.

Die gefundenen erfindungsgemäßen Nukleinsäure-Sequenzen können auch genomische oder mRNA-Sequenzen sein.

Die Erfindung betrifft auch genomische Gene, ihre Exon- und Intronstruktur und deren Spleißvarianten, erhältlich aus den cDNAs der Sequenzen Seq. ID No. 1-50, sowie deren Verwendung zusammen mit geeigneten regulativen Elementen, wie geeigneten Promotoren und/oder Enhancern. 40

Mit den erfindungsgemäßen Nukleinsäuren (cDNA-Sequenzen) Seq. ID No. 1-50 werden genomische BAC-, PAC- und Cosmid-Bibliotheken gescreent und über komplementäre Basenpaarung (Hybridisierung) spezifisch humane Klone isoliert. Die so isolierten BAC-, PAC- und Cosmid-Klone werden mit Hilfe der Fluoreszenz-in-situ-Hybridisation auf Metaphasenchromosomen hybridisiert und entsprechende Chromosomenabschnitte identifiziert, auf denen die entsprechenden genomischen Gene liegen. BAC-, PAC- und Cosmid-Klone werden sequenziert, um die entsprechenden genomischen Gene in ihrer vollständigen Struktur (Promotoren, Enhancer, Silencer, Exons und Introns) aufzuklären. BAC-, PAC- und Cosmid-Klone können als eigenständige Moleküle für den Gentransfer eingesetzt werden (s. Fig. 5). 45

Die Erfindung betrifft auch BAC-, PAC- und Cosmid-Klone, enthaltend funktionelle Gene und ihre chromosomale Lokalisation, entsprechend den Sequenzen Seq. ID. No. 1 bis Seq. ID No. 50, zur Verwendung als Vehikel zum Gentransfer. 50

Bedeutungen von Fachbegriffen und Abkürzungen

Nukleinsäuren = Unter Nukleinsäuren sind in der vorliegenden Erfindung zu verstehen: mRNA, partielle cDNA, vollständigen cDNA und genomische Gene (Chromosomen). 55

ORF = Open Reading Frame, eine definierte Abfolge von Aminosäuren, die von der cDNA-Sequenz abgeleitet werden kann.

Contig = eine Menge von DNA-Sequenzen, die aufgrund sehr großer Ähnlichkeiten zu einer Sequenz zusammengefaßt werden können (Consensus) 60

Singleton = ein Contig, der nur eine Sequenz enthält.

Erklärung zu den Alignmentparametern

minimal initial match = minimaler anfänglicher Identitätsbereich 65

maximum pads per read = maximale Anzahl von Insertionen

maximum percent mismatch = maximale Abweichung in %.

Erklärung der Abbildungen

Fig. 1 zeigt die systematische Gen-Suche in der Incyte LifeSeq Datenbank.

Fig. 2a zeigt das Prinzip der EST-Assemblierung

Fig. 2b1-2b4 zeigt das gesamte Prinzip der EST-Assemblierung

Fig. 3 zeigt die in silico Subtraktion der Genexpression in verschiedenen Geweben

Fig. 4a zeigt die Bestimmung der gewebsspezifischen Expression über elektronischen Northern.

Fig. 4b zeigt den elektronischen Northern

Fig. 5 zeigt die Isolierung von genomischen BAC- und PAC-Klonen.

Die nachfolgenden Beispiele erläutern die Herstellung der erfindungsgemäßen Nukleinsäure-Sequenzen, ohne die Erfindung auf diese Beispiele und Nukleinsäure-Sequenzen zu beschränken.

Beispiel 1

Suche nach Tumor-bezogenen Kandidatengenen

Zuerst wurden sämtliche ESTs des entsprechenden Gewebes aus der LifeSeq-Datenbank (vom Oktober 1997) extrahiert. Diese wurden dann mittels des Programms GAP4 des Staden-Pakets mit den Parametern 0% mismatch, 8 pads per read und einem minimalen match von 20 assembliert. Die nicht in die GAP4-Datenbank aufgenommenen Sequenzen (Fails) wurden erst bei 1% mismatch und dann nochmals bei 2% mismatch mit der Datenbank assembliert. Aus den Contigs der Datenbank, die aus mehr als einer Sequenz bestanden, wurden Consensussequenzen errechnet. Die Singletons der Datenbank, die nur aus einer Sequenz bestanden, wurden mit den nicht in die GAP4-Datenbank aufgenommenen Sequenzen bei 2% mismatch erneut assembliert. Wiederum wurden für die Contigs die Consensussequenzen ermittelt. Alle übrigen ESTs wurden bei 4% mismatch erneut assembliert. Die Consensussequenzen wurden abermals extrahiert und mit den vorherigen Consensussequenzen sowie den Singletons und den nicht in die Datenbank aufgenommenen Sequenzen abschließend bei 4% mismatch assembliert. Die Consensussequenzen wurden gebildet und mit den Singletons und Fails als Ausgangsbasis für die Gewebsvergleiche verwendet. Durch diese Prozedur konnte sichergestellt werden, daß unter den verwendeten Parametern sämtliche Sequenzen von einander unabhängige Genbereiche darstellten.

Fig. 2b1-2b4 veranschaulicht die Verlängerung der Blasengewebe ESTs.

Die so assemblierten Sequenzen der jeweiligen Gewebe wurden anschließend mittels des gleichen Programms miteinander verglichen (Fig. 3). Hierzu wurden erst alle Sequenzen des ersten Gewebes in die Datenbank eingegeben. (Daher war es wichtig, daß diese voneinander unabhängig waren.)

Dann wurden alle Sequenzen des zweiten Gewebes mit allen des ersten verglichen. Das Ergebnis waren Sequenzen, die für das erste bzw. das zweite Gewebe spezifisch waren, sowie welche, die in beiden vorkamen. Bei Letzteren wurde das Verhältnis der Häufigkeit des Vorkommens in den jeweiligen Geweben ausgewertet. Sämtliche, die Auswertung der assemblierten Sequenzen betreffenden Programme, wurden selbst entwickelt.

Alle Sequenzen, die mehr als viermal in jeweils einem der verglichenen Gewebe vorkamen, sowie alle, die mindestens fünfmal so häufig in einem der beiden Gewebe vorkamen wurden weiter untersucht. Diese Sequenzen wurden einem elektronischen Northern (s. Beispiel 2.1) unterzogen, wodurch die Verteilung in sämtlichen Tumor- und Normal-Geweben untersucht wurde (s. Fig. 4a und Fig. 4b). Die relevanten Kandidaten wurden dann mit Hilfe sämtlicher Incyte ESTs und allen ESTs öffentlicher Datenbanken verlängert (s. Beispiel 3). Anschließend wurden die Sequenzen und ihre Übersetzung in mögliche Proteine mit allen Nukleotid- und Proteindatenbanken verglichen, sowie auf mögliche, für Proteine kodierende Regionen untersucht.

Beispiel 2

Algorithmus zur Identifikation und Verlängerung von partiellen cDNA-Sequenzen mit verändertem Expressionsmuster

Im folgenden soll ein Algorithmus zur Auffindung über- oder unterexprimierter Gene erläutert werden. Die einzelnen Schritte sind der besseren Übersicht halber auch in einem Flußdiagramm zusammengefaßt (s. Fig. 4b).

2.1 Elektronischer Northern-Blot

Zu einer partiellen DNA-Sequenz S, z. B. einem einzelnen EST oder einem Contig von ESTs, werden mittels eines Standardprogramms zur Homologiesuche, z. B. BLAST (Altschul, S. F., Gish W., Miller, W., Myers, E. W. und Lipman, D. J. (1990) J. Mol. Biol., 215, 403-410), BLAST2 (Altschul, S. F., Madden, T. L., Schäffer, A. A., Zhang, J., Zhang, Z., Miller, W. und Lipman, D. J. (1997) Nucleic Acids Research 25 3389-3402) oder FASTA (Pearson, W. R. und Lipman, D. J. (1988) Proc. Natl. Acad. Sci. USA 85 2444-2448), die homologen Sequenzen in verschiedenen nach Geweben geordneten (privaten oder öffentlichen) EST-Bibliotheken bestimmt. Die dadurch ermittelten (relativen oder absoluten) Gewebe-spezifischen Vorkommenshäufigkeiten dieser Partial-Sequenz S werden als elektronischer Northern-Blot bezeichnet.

2.1.1

Analog der unter 2.1 beschriebenen Verfahrensweise wurde die Sequenz Seq. ID No. 16 gefunden, die 17,7.x stärker im normalen Blasentumorgewebe als im normalem Blasengewebe vorkommt.

Das Ergebnis ist wie folgt:

	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0039	0.0690	0.0565	17.6998	5
Brust	0.0000	0.0000	undef	undef	
Duennndarm	0.0031	0.0000	undef	0.0000	
Eierstock	0.0180	0.0078	2.3025	0.4343	
Endokrines_Gewebe	0.0000	0.0000	undef	undef	
Gastrointestinal	0.0000	0.0000	undef	undef	
Gehirn	0.0000	0.0000	undef	undef	10
Haematopoetisch	0.0000	0.0000	undef	undef	
Haut	0.0000	0.0000	undef	undef	
Hepatisch	0.0000	0.0000	undef	undef	
Herz	0.0000	0.0000	undef	undef	
Hoden	0.0000	0.0000	undef	undef	
Lunge	0.0000	0.0000	undef	undef	15
Magen-Speiserohre	0.0000	0.0000	undef	undef	
Muskel-Skelett	0.0000	0.0000	undef	undef	
Niere	0.0000	0.0000	undef	undef	
Pankreas	0.0000	0.0000	undef	undef	
Penis	0.0000	0.0000	undef	undef	
Prostata	0.0065	0.0064	1.0236	0.9769	20
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0000	undef	undef	
Uterus_allgemein	0.0000	0.0000	undef	undef	
Brust-Hyperplasie	0.0000				
Prostata-Hyperplasie	0.0178				25
Samenblase	0.0000				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0000				
Zervix	0.0000				30
FOETUS					
	%Haeufigkeit				
Entwicklung	0.0000				
Gastrointestinal	0.0028				
Gehirn	0.0000				35
Haematopoetisch	0.0000				
Haut	0.0000				
Hepatisch	0.0000				
Herz-Blutgefuesse	0.0000				
Lunge	0.0000				
Nebenniere	0.0000				40
Niere	0.0000				
Placenta	0.0000				
Prostata	0.0000				
Sinnesorgane	0.0000				45
NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN					
	%Haeufigkeit				
Brust	0.0000				
Eierstock_n	0.0000				
Eierstock_t	0.0000				
Endokrines_Gewebe	0.0000				50
Foetal	0.0035				
Gastrointestinal	0.0000				
Haematopoetisch	0.0000				
Haut-Muskel	0.0000				
Hoden	0.0000				55
Lunge	0.0000				
Nerven	0.0000				
Prostata	0.0068				
Sinnesorgane	0.0000				
Uterus_n	0.0000				60

In analoger Verfahrensweise wurden auch folgende Northern gefunden:

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0281	0.0000	undef
	Brust	0.0038	0.0056	0.6805	1.4694
	Duendarm	0.0399	0.0000	undef	0.0000
	Eierstock	0.0000	0.0078	0.0000	undef
	Endokrines_Gewebe	0.0000	0.0000	undef	undef
	Gastrointestinal	0.0479	0.0000	undef	0.0000
10	Gehirn	0.0000	0.0010	0.0000	undef
	Haematopoetisch	0.0227	0.0000	undef	0.0000
	Haut	0.0037	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0323	0.0000	undef
	Herz	0.0000	0.0000	undef	undef
	Hoden	0.0000	0.0585	0.0000	undef
15	Lunge	0.0145	0.0123	1.1854	0.8436
	Magen-Speiserohre	0.0000	0.0000	undef	undef
	Muskel-Skelett	0.0120	0.0120	0.9994	1.0006
	Niere	0.0000	0.0274	0.0000	undef
	Pankreas	0.0066	0.0110	0.5983	1.6714
20	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0022	0.0021	1.0236	0.9769
	Uterus_Endometrium	0.0000	0.0000	undef	undef
	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0128			
25	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0000			
	Sinnesorgane	0.0000			
	Weisse_Blutkoerperchen	0.0009			
	Zervix	0.0000			

30		FOETUS
		%Haeufigkeit
	Entwicklung	0.0000
	Gastrointestinal	0.0000
35	Gehirn	0.0000
	Haematopoetisch	0.0039
	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
40	Nebenniere	0.0000
	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

45		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
	Brust	0.0000
	Eierstock_n	0.0000
50	Eierstock_t	0.0000
	Endokrines_Gewebe	0.0000
	Foetal	0.0000
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
	Haut-Muskel	0.0000
55	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000
60	Uterus_n	0.0000

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Elektronischer Northern für SEQ. ID. NO: 2

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0000	0.0307	0.0000 undef	5
Brust	0.0307	0.0376	0.8166 1.2245	
Duennndarm	0.0337	0.0165	2.0391 0.4904	
Eierstock	0.0120	0.0364	0.3289 3.0402	
Endokrines_Gewebe	0.0255	0.0075	3.3962 0.2944	
Gastrointestinal	0.0153	0.0185	0.8283 1.2072	10
Gehirn	0.0185	0.0216	0.8571 1.1667	
Haematopoetisch	0.0201	0.0379	0.5293 1.8892	
Haut	0.0844	0.0000	undef 0.0000	
Hepatisch	0.0238	0.0065	3.6765 0.2720	
Herz	0.0148	0.0000	undef 0.0000	15
Hoden	0.0575	0.0351	1.6399 0.6098	
Lunge	0.0145	0.0082	1.7781 0.5624	
Magen-Speiserohre	0.0387	0.0077	5.0421 0.1983	
Muskel-Skelett	0.0308	0.0300	1.0280 0.9728	
Niere	0.0217	0.0000	undef 0.0000	20
Pankreas	0.0099	0.0110	0.8974 1.1143	
Penis	0.0240	0.0000	undef 0.0000	
Prostata	0.0262	0.0213	1.2284 0.8141	
Uterus_Endometrium	0.0135	0.0000	undef 0.0000	
Uterus_Myometrium	0.0152	0.0408	0.3741 2.6732	25
Uterus_allgemein	0.0204	0.0954	0.2135 4.6839	
Brust-Hyperplasie	0.0512			
Prostata-Hyperplasie	0.0268			
Samenblase	0.0089			
Sinnesorgane	0.0235			
Weisse_Blutkoerperchen	0.0286			30
Zervix	0.0106			
FOETUS				
	%Haeufigkeit			35
Entwicklung	0.0278			
Gastrointestinal	0.0305			
Gehirn	0.0063			
Haematopoetisch	0.0157			
Haut	0.0000			40
Hepatisch	0.0000			
Herz-Blutgefuesse	0.0213			
Lunge	0.0289			
Nebenniere	0.0000			
Niere	0.0185			45
Placenta	0.0121			
Prostata	0.0000			
Sinnesorgane	0.0126			
NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN				
	%Haeufigkeit			50
Brust	0.0204			
Eierstock_n	0.0000			
Eierstock_t	0.0051			55
Endokrines_Gewebe	0.0000			
Foetal	0.0122			
Gastrointestinal	0.0488			
Haematopoetisch	0.0000			
Haut-Muskel	0.0000			60
Hoden	0.0463			
Lunge	0.0164			
Nerven	0.0100			
Prostata	0.0137			
Sinnesorgane	0.0000			
Uterus_n	0.0125			65

		NORMAL		TUMOR		Verhaeltnisse	
		%Haeufigkeit		%Haeufigkeit		N/T	T/N
5	Blase	0.0000	0.0256	0.0000	undef	undef	undef
	Brust	0.0000	0.0000	undef	undef	undef	undef
	Duendarm	0.0000	0.0000	undef	undef	undef	undef
	Eierstock	0.0000	0.0000	undef	undef	undef	undef
	Endokrines_Gewebe	0.0000	0.0000	undef	undef	undef	undef
10	Gastrointestinal	0.0000	0.0000	undef	undef	undef	undef
	Gehirn	0.0000	0.0000	undef	undef	undef	undef
	Haematopoetisch	0.0000	0.0000	undef	undef	undef	undef
	Haut	0.0000	0.0000	undef	undef	undef	undef
	Hepatisch	0.0000	0.0000	undef	undef	undef	undef
15	Herz	0.0000	0.0000	undef	undef	undef	undef
	Hoden	0.0000	0.0000	undef	undef	undef	undef
	Lunge	0.0000	0.0000	undef	undef	undef	undef
	Magen-Speiserohre	0.0000	0.0000	undef	undef	undef	undef
	Muskel-Skelett	0.0000	0.0000	undef	undef	undef	undef
20	Niere	0.0000	0.0000	undef	undef	undef	undef
	Pankreas	0.0000	0.0000	undef	undef	undef	undef
	Penis	0.0030	0.0000	undef	0.0000	undef	undef
	Prostata	0.0000	0.0000	undef	undef	undef	undef
	Uterus_Endometrium	0.0000	0.0000	undef	undef	undef	undef
	Uterus_Myometrium	0.0000	0.0000	undef	undef	undef	undef
25	Uterus_allgemein	0.0000	0.0000	undef	undef	undef	undef
	Brust-Hyperplasie	0.0000					
	Prostata-Hyperplasie	0.0000					
	Samenblase	0.0000					
	Sinnesorgane	0.0000					
30	Weisse_Blutkoerperchen	0.0000					
	Zervix	0.0000					

		FOETUS	
		%Haeufigkeit	
35	Entwicklung	0.0000	
	Gastrointestinal	0.0000	
	Gehirn	0.0000	
	Haematopoetisch	0.0000	
40	Haut	0.0000	
	Hepatisch	0.0000	
	Herz-Blutgefuesse	0.0000	
	Lunge	0.0000	
	Nebenniere	0.0000	
45	Niere	0.0000	
	Placenta	0.0000	
	Prostata	0.0000	
	Sinnesorgane	0.0000	

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
		%Haeufigkeit	
		Brust	0.0000
		Eierstock_n	0.0000
55	Eierstock_t	0.0000	
	Endokrines_Gewebe	0.0000	
	Foetal	0.0000	
	Gastrointestinal	0.0000	
	Haematopoetisch	0.0000	
60	Haut-Muskel	0.0000	
	Hoden	0.0000	
	Lunge	0.0000	
	Nerven	0.0000	
	Prostata	0.0000	
65	Sinnesorgane	0.0000	
	Uterus_n	0.0000	

	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0000	0.0230	0.0000	undef	5
Brust	0.0090	0.0094	0.9527	1.0496	
Duenn darm	0.0123	0.0165	0.7415	1.3487	
Eierstock	0.0150	0.0104	1.4391	0.6949	
Endokrines Gewebe	0.0068	0.0125	0.5434	1.8403	
Gastrointestinal	0.0038	0.0046	0.8283	1.2072	10
Gehirn	0.0044	0.0123	0.3600	2.7779	
Haematopoetisch	0.0187	0.0000	undef	0.0000	
Haut	0.0000	0.0000	undef	undef	
Hepatisch	0.0000	0.0388	0.0000	undef	
Herz	0.0138	0.0000	undef	0.0000	15
Hoden	0.0173	0.0000	undef	0.0000	
Lunge	0.0093	0.0143	0.6532	1.5310	
Magen-Speiserohre	0.0097	0.0000	undef	0.0000	
Muskel-Skelett	0.0017	0.0180	0.0952	10.5060	
Niere	0.0136	0.0137	0.9913	1.0088	20
Pankreas	0.0149	0.0055	2.6923	0.3714	
Penis	0.0150	0.0000	undef	0.0000	
Prostata	0.0065	0.0064	1.0236	0.9769	
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0136	0.0000	undef	25
Uterus_allgemein	0.0000	0.0000	undef	undef	
Brust-Hyperplasie	0.0032				
Prostata-Hyperplasie	0.0178				
Samenblase	0.0089				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0035				30
Zervix	0.0000				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0167	
Gehirn	0.0000	
Haematopoetisch	0.0354	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0072	
Nebenniere	0.0000	
Niere	0.0062	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0272	
Eierstock_n	0.0000	
Eierstock_t	0.0051	55
Endokrines_Gewebe	0.0000	
Foetal	0.0151	
Gastrointestinal	0.0000	
Haematopoetisch	0.0114	
Haut-Muskel	0.0259	
Hoden	0.0077	60
Lunge	0.0000	
Nerven	0.0060	
Prostata	0.0068	
Sinnesorgane	0.0155	
Uterus_n	0.0250	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0204	0.0000	undef
	Brust	0.0026	0.0056	0.4537	2.2042
	Duennndarm	0.0031	0.0000	undef	0.0000
	Eierstock	0.0060	0.0156	0.3838	2.6058
	Endokrines_Gewebe	0.0051	0.0000	undef	0.0000
10	Gastrointestinal	0.0134	0.0093	1.4496	0.6898
	Gehirn	0.0052	0.0051	1.0079	0.9921
	Haematopoetisch	0.0000	0.0000	undef	undef
	Haut	0.0000	0.0000	undef	undef
	Hepatisch	0.0000	0.0000	undef	undef
15	Herz	0.0032	0.0000	undef	0.0000
	Hoden	0.0000	0.0000	undef	undef
	Lunge	0.0031	0.0020	1.5241	0.6561
	Magen-Speiseroehre	0.0000	0.0000	undef	undef
	Muskel-Skelett	0.0034	0.0060	0.5711	1.7510
20	Niere	0.0190	0.0068	2.7756	0.3603
	Pankreas	0.0017	0.0000	undef	0.0000
	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0131	0.0043	3.0709	0.3256
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0059			
	Samenblase	0.0000			
	Sinnesorgane	0.0000			
30	Weisse_Blutkoerperchen	0.0000			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0083
	Gehirn	0.0063
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHLIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.0000
55	Eierstock_t	0.0051
	Endokrines_Gewebe	0.0000
	Foetal	0.0000
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0082
	Nerven	0.0050
	Prostata	0.0137
65	Sinnesorgane	0.0000
	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0000	0.0204	0.0000 undef	5
Brust	0.0038	0.0075	0.5104 1.9593	
Duenn darm	0.0031	0.0000	undef 0.0000	
Eierstock	0.0060	0.0052	1.1513 0.8686	
Endokrines Gewebe	0.0017	0.0000	undef 0.0000	
Gastrointestinal	0.0038	0.0000	undef 0.0000	10
Gehirn	0.0022	0.0021	1.0799 0.9260	
Haematopoetisch	0.0053	0.0000	undef 0.0000	
Haut	0.0000	0.0000	undef undef	
Hepatisch	0.0048	0.0000	undef 0.0000	
Herz	0.0042	0.0137	0.3084 3.2426	15
Hoden	0.0000	0.0000	undef undef	
Lunge	0.0052	0.0143	0.3629 2.7557	
Magen-Speiserohre	0.0000	0.0000	undef undef	
Muskel-Skelett	0.0017	0.0060	0.2856 3.5020	
Niere	0.0000	0.0000	undef undef	20
Pankreas	0.0017	0.0110	0.1496 6.6857	
Penis	0.0000	0.0000	undef undef	
Prostata	0.0000	0.0021	0.0000 undef	
Uterus_Endometrium	0.0135	0.0000	undef 0.0000	
Uterus_Myometrium	0.0229	0.0068	3.3668 0.2970	25
Uterus_allgemein	0.0204	0.0000	undef 0.0000	
Brust-Hyperplasie	0.0000			
Prostata-Hyperplasie	0.0000			
Samenblase	0.0000			
Sinnesorgane	0.0353			
Weisse_Blutkoerperchen	0.0009			30
Zervix	0.0000			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0000	
Haematopoetisch	0.0118	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0036	
Nebenniere	0.0000	
Niere	0.0124	45
Placenta	0.0242	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0000	50
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0245	55
Foetal	0.0151	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0259	60
Hoden	0.0000	
Lunge	0.0164	
Nerven	0.0080	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0250	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0179	0.0000	undef
	Brust	0.0038	0.0132	0.2917	3.4287
	Duendarm	0.0061	0.0000	undef	0.0000
	Eierstock	0.0060	0.0052	1.1513	0.8686
	Endokrines Gewebe	0.0017	0.0100	0.1698	5.8889
10	Gastrointestinal	0.0019	0.0370	0.0518	19.3158
	Gehirn	0.0096	0.0051	1.8719	0.5342
	Haematopoetisch	0.0067	0.0000	undef	0.0000
	Haut	0.0000	0.0000	undef	undef
	Hepatisch	0.0048	0.0065	0.7353	1.3600
15	Herz	0.0138	0.0137	1.0023	0.9977
	Hoden	0.0288	0.0468	0.6150	1.6261
	Lunge	0.0031	0.0143	0.2177	4.5929
	Magen-Speiserohre	0.0387	0.0153	2.5211	0.3967
	Muskel-Skelett	0.0017	0.0000	undef	0.0000
20	Niere	0.0054	0.0000	undef	0.0000
	Pankreas	0.0050	0.0055	0.8974	1.1143
	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0000	0.0021	0.0000	undef
	Uterus_Endometrium	0.0000	0.0000	undef	undef
25	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0102	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0000			
	Samenblase	0.0000			
	Sinnesorgane	0.0000			
30	Weisse_Blutkoerperchen	0.0026			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0028
	Gehirn	0.0751
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0036
	Lunge	0.0036
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0061
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0068
	Eierstock_n	0.0000
55	Eierstock_t	0.0000
	Endokrines Gewebe	0.0000
	Foetal	0.0087
	Gastrointestinal	0.0122
	Haematopoetisch	0.0114
60	Haut-Muskel	0.0097
	Hoden	0.0540
	Lunge	0.0082
	Nerven	0.0201
	Prostata	0.0205
65	Sinnesorgane	0.0000
	Uterus_n	0.0375

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0000	0.0179	0.0000	undef	5
Brust	0.0013	0.0038	0.3403	2.9389	
Duennndarm	0.0123	0.0000	undef	0.0000	
Eierstock	0.0060	0.0078	0.7675	1.3029	
Endokrines_Gewebe	0.0068	0.0100	0.6792	1.4722	
Gastrointestinal	0.0038	0.0093	0.4142	2.4145	10
Gehirn	0.0044	0.0144	0.3086	3.2409	
Haematopoetisch	0.0080	0.0000	undef	0.0000	
Haut	0.0037	0.0000	undef	0.0000	
Hepatisch	0.0000	0.0000	undef	undef	
Herz	0.0021	0.0000	undef	0.0000	
Hoden	0.0058	0.0000	undef	0.0000	15
Lunge	0.0031	0.0082	0.3810	2.6245	
Magen-Speiserohre	0.0000	0.0307	0.0000	undef	
Muskel-Skelett	0.0051	0.0060	0.8567	1.1673	
Niere	0.0000	0.0000	undef	undef	
Pankreas	0.0066	0.0000	undef	0.0000	20
Penis	0.0060	0.0000	undef	0.0000	
Prostata	0.0087	0.0064	1.3648	0.7327	
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0068	0.0000	undef	
Uterus_allgemein	0.0051	0.0000	undef	0.0000	25
Brust-Hyperplasie	0.0000				
Prostata-Hyperplasie	0.0059				
Samenblase	0.0000				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0035				30
Zervix	0.0000				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0056	
Gehirn	0.0000	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0036	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0061	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0204	
Eierstock_n	0.0000	
Eierstock_t	0.0608	
Endokrines_Gewebe	0.0000	55
Foetal	0.0029	
Gastrointestinal	0.0244	
Haematopoetisch	0.0000	
Haut-Muskel	0.0097	
Hoden	0.0154	60
Lunge	0.0000	
Nerven	0.0080	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0179	0.0000	undef
	Brust	0.0064	0.0056	1.1342	0.8817
	Duennndarm	0.0000	0.0165	0.0000	undef
	Eierstock	0.0030	0.0026	1.1513	0.8686
	Endokrines_Gewebe	0.0153	0.0025	6.1132	0.1636
10	Gastrointestinal	0.0038	0.0046	0.8283	1.2072
	Gehirn	0.0015	0.0051	0.2880	3.4724
	Haematopoetisch	0.0067	0.0000	undef	0.0000
	Haut	0.0037	0.0847	0.0433	23.0839
	Hepatisch	0.0000	0.0065	0.0000	undef
15	Herz	0.0053	0.0000	undef	0.0000
	Hoden	0.0230	0.0234	0.9839	1.0163
	Lunge	0.0042	0.0041	1.0161	0.9842
	Magen-Speiserohre	0.0000	0.0077	0.0000	undef
	Muskel-Skelett	0.0034	0.0000	undef	0.0000
20	Niere	0.0054	0.0068	0.7930	1.2610
	Pankreas	0.0017	0.0000	undef	0.0000
	Penis	0.0030	0.0267	0.1123	8.9035
	Prostata	0.0065	0.0064	1.0236	0.9769
	Uterus_Endometrium	0.0135	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0152	0.0000	undef	0.0000
	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0032			
	Prostata-Hyperplasie	0.0000			
	Samenblase	0.0000			
30	Sinnesorgane	0.0000			
	Weisse_Blutkoerperchen	0.0043			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0039
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0036
	Lunge	0.0108
	Nebenniere	0.0000
45	Niere	0.0062
	Placenta	0.0000
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0068
	Eierstock_n	0.0000
55	Eierstock_t	0.0101
	Endokrines_Gewebe	0.0000
	Foetal	0.0047
	Gastrointestinal	0.0000
	Haematopoetisch	0.0114
60	Haut-Muskel	0.0194
	Hoden	0.0309
	Lunge	0.0000
	Nerven	0.0040
	Prostata	0.0000
65	Sinnesorgane	0.0000
	Uterus_n	0.0125

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0000	0.0153	0.0000	undef	5
Brust	0.0051	0.0019	2.7221	0.3674	
Duennndarm	0.0000	0.0000	undef	undef	
Eierstock	0.0000	0.0000	undef	undef	10
Endokrines_Gewebe	0.0000	0.0000	undef	undef	
Gastrointestinal	0.0000	0.0093	0.0000	undef	
Gehirn	0.0000	0.0113	0.0000	undef	
Haematopoetisch	0.0000	0.0000	undef	undef	15
Haut	0.0000	0.0000	undef	undef	
Hepatisch	0.0000	0.0129	0.0000	undef	
Herz	0.0011	0.0000	undef	0.0000	
Hoden	0.0058	0.0000	undef	0.0000	
Lunge	0.0021	0.0000	undef	0.0000	20
Magen-Speiserohre	0.0000	0.0000	undef	undef	
Muskel-Skelett	0.0000	0.0000	undef	undef	
Niere	0.0000	0.0000	undef	undef	
Pankreas	0.0000	0.0000	undef	undef	25
Penis	0.0000	0.0000	undef	undef	
Prostata	0.0000	0.0000	undef	undef	
Uterus_Endometrium	0.0000	0.0000	undef	undef	30
Uterus_Myometrium	0.0000	0.0000	undef	undef	
Uterus_allgemein	0.0000	0.0000	undef	undef	
Brust-Hyperplasie	0.0000				
Prostata-Hyperplasie	0.0000				35
Samenblase	0.0000				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0000				
Zervix	0.0000				

FOETUS					
	%Haeufigkeit				
Entwicklung	0.0000				40
Gastrointestinal	0.0000				
Gehirn	0.0000				
Haematopoetisch	0.0000				45
Haut	0.0000				
Hepatisch	0.0000				
Herz-Blutgefuesse	0.0036				50
Lunge	0.0000				
Nebenniere	0.0000				
Niere	0.0000				55
Placenta	0.0000				
Prostata	0.0000				
Sinnesorgane	0.0000				

NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN					
	%Haeufigkeit				
Brust	0.0000				60
Eierstock_n	0.0000				
Eierstock_t	0.0000				
Endokrines_Gewebe	0.0000				65
Foetal	0.0006				
Gastrointestinal	0.0000				
Haematopoetisch	0.0000				70
Haut-Muskel	0.0000				
Hoden	0.0000				
Lunge	0.0000				75
Nerven	0.0010				
Prostata	0.0000				
Sinnesorgane	0.0000				80
Uterus_n	0.0000				

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0153	0.0000	undef
	Brust	0.0038	0.0038	1.0208	0.9796
	Duennndarm	0.0000	0.0000	undef	undef
	Eierstock	0.0000	0.0104	0.0000	undef
	Endokrines_Gewebe	0.0068	0.0025	2.7170	0.3681
10	Gastrointestinal	0.0077	0.0093	0.8283	1.2072
	Gehirn	0.0000	0.0062	0.0000	undef
	Haematopoetisch	0.0107	0.0000	undef	0.0000
	Haut	0.0073	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0194	0.0000	undef
15	Herz	0.0021	0.0000	undef	0.0000
	Hoden	0.0000	0.0000	undef	undef
	Lunge	0.0010	0.0041	0.2540	3.9367
	Magen-Speiserohre	0.0097	0.0077	1.2605	0.7933
	Muskel-Skelett	0.0000	0.0000	undef	undef
20	Niere	0.0054	0.0068	0.7930	1.2610
	Pankreas	0.0050	0.0000	undef	0.0000
	Penis	0.0000	0.0267	0.0000	undef
	Prostata	0.0065	0.0298	0.2193	4.5590
	Uterus_Endometrium	0.0068	0.0528	0.1280	7.8106
25	Uterus_Myometrium	0.0076	0.0068	1.1223	0.8911
	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0238			
	Samenblase	0.0000			
	Sinnesorgane	0.0000			
30	Weisse_Blutkoerperchen	0.0026			
	Zervix	0.0106			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0139
	Gastrointestinal	0.0194
	Gehirn	0.0063
	Haematopoetisch	0.0000
40	Haut	0.0000
	Hepatisch	0.0260
	Herz-Blutgefuesse	0.0071
	Lunge	0.0108
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0121
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.0000
55	Eierstock_t	0.0557
	Endokrines_Gewebe	0.0000
	Foetal	0.0076
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0030
	Prostata	0.0137
65	Sinnesorgane	0.0387
	Uterus_n	0.0042

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0000	0.0153	0.0000 undef	5
Brust	0.0013	0.0000	undef 0.0000	
Duenn darm	0.0031	0.0000	undef 0.0000	
Eierstock	0.0000	0.0026	0.0000 undef	
Endokrines Gewebe	0.0017	0.0075	0.2264 4.4166	
Gastrointestinal	0.0000	0.0093	0.0000 undef	10
Gehirn	0.0044	0.0000	undef 0.0000	
Haematopoetisch	0.0000	0.0000	undef undef	
Haut	0.0037	0.0000	undef 0.0000	
Hepatisch	0.0000	0.0000	undef undef	
Herz	0.0000	0.0000	undef undef	
Hoden	0.0000	0.0117	0.0000 undef	15
Lunge	0.0000	0.0000	undef undef	
Magen-Speiserohre	0.0000	0.0000	undef undef	
Muskel-Skelett	0.0000	0.0000	undef undef	
Niere	0.0000	0.0000	undef undef	
Pankreas	0.0017	0.0000	undef 0.0000	20
Penis	0.0000	0.0000	undef undef	
Prostata	0.0000	0.0021	0.0000 undef	
Uterus_Endometrium	0.0000	0.0000	undef undef	
Uterus_Myometrium	0.0000	0.0000	undef undef	
Uterus_allgemein	0.0000	0.0000	undef undef	25
Brust-Hyperplasie	0.0032			
Prostata-Hyperplasie	0.0000			
Samenblase	0.0000			
Sinnesorgane	0.0000			
Weisse_Blutkoerperchen	0.0000			30
Zervix	0.0000			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0125	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0036	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0000	50
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0245	55
Foetal	0.0023	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	60
Hoden	0.0000	
Lunge	0.0082	
Nerven	0.0020	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0000	0.0153	0.0000	undef
	Brust	0.0090	0.0169	0.5293	1.8893
	Duenn darm	0.0031	0.0000	undef	0.0000
	Eierstock	0.0060	0.0052	1.1513	0.8686
	Endokrines Gewebe	0.0051	0.0226	0.2264	4.4166
10	Gastrointestinal	0.0057	0.0231	0.2485	4.0241
	Gehirn	0.0052	0.0082	0.6300	1.5874
	Haematopoetisch	0.0107	0.0000	undef	0.0000
	Haut	0.0037	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0000	undef	undef
15	Herz	0.0138	0.0000	undef	0.0000
	Hoden	0.0345	0.0117	2.9518	0.3388
	Lunge	0.0021	0.0123	0.1693	5.9051
	Magen-Speiserohre	0.0000	0.0000	undef	undef
	Muskel-Skelett	0.0051	0.0060	0.8567	1.1673
20	Niere	0.0163	0.0068	2.3791	0.4203
	Pankreas	0.0000	0.0055	0.0000	undef
	Penis	0.0030	0.0267	0.1123	8.9035
	Prostata	0.0174	0.0128	1.3648	0.7327
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0051	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0149			
	Samenblase	0.0267			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0087			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0139
	Gastrointestinal	0.0139
	Gehirn	0.0063
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0107
	Lunge	0.0072
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0061
	Prostata	0.0249
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.0000
55	Eierstock_t	0.0101
	Endokrines_Gewebe	0.0000
	Foetal	0.0146
	Gastrointestinal	0.0000
	Haematopoetisch	0.0057
60	Haut-Muskel	0.0000
	Hoden	0.0309
	Lunge	0.0000
	Nerven	0.0010
	Prostata	0.0137
65	Sinnesorgane	0.0000
	Uterus_n	0.0083

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0000	0.0153	0.0000 undef	5
Brust	0.0051	0.0395	0.1296 7.7146	
Duendarm	0.0000	0.0000	undef undef	
Eierstock	0.0060	0.0234	0.2558 3.9088	
Endokrines_Gewebe	0.0000	0.0000	undef undef	
Gastrointestinal	0.0115	0.0000	undef 0.0000	10
Gehirn	0.0000	0.0041	0.0000 undef	
Haematopoetisch	0.0027	0.0000	undef 0.0000	
Haut	0.0037	0.0000	undef 0.0000	
Hepatisch	0.0381	0.0129	2.9412 0.3400	
Herz	0.0021	0.0000	undef 0.0000	
Hoden	0.0000	0.0234	0.0000 undef	15
Lunge	0.0073	0.0245	0.2964 3.3743	
Magen-Speiserohre	0.0000	0.0537	0.0000 undef	
Muskel-Skelett	0.0137	0.0420	0.3263 3.0643	
Niere	0.0163	0.0000	undef 0.0000	
Pankreas	0.0017	0.0331	0.0499 20.0570	20
Penis	0.0000	0.0267	0.0000 undef	
Prostata	0.0022	0.0128	0.1706 5.8615	
Uterus_Endometrium	0.0068	0.0000	undef 0.0000	
Uterus_Myometrium	0.0305	0.0136	2.2445 0.4455	
Uterus_allgemein	0.0000	0.0000	undef undef	25
Brust-Hyperplasie	0.0000			
Prostata-Hyperplasie	0.0000			
Samenblase	0.0000			
Sinnesorgane	0.0118			
Weisse_Blutkoerperchen	0.0000			30
Zervix	0.0319			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0278	35
Gastrointestinal	0.0583	
Gehirn	0.0000	
Haematopoetisch	0.0354	
Haut	0.5025	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0071	
Lunge	0.0542	
Nebenniere	0.0761	
Niere	0.1235	45
Placenta	0.0727	
Prostata	0.0249	
Sinnesorgane	0.1255	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0204	
Eierstock_n	0.0000	
Eierstock_t	0.0709	55
Endokrines_Gewebe	0.0245	
Foetal	0.0175	
Gastrointestinal	0.0122	
Haematopoetisch	0.0000	
Haut-Muskel	0.0162	60
Hoden	0.0000	
Lunge	0.0000	
Nerven	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0039	0.1278	0.0305	32.7774
	Brust	0.0179	0.0414	0.4331	2.3091
	Duenn darm	0.0031	0.1323	0.0232	43.1571
	Eierstock	0.0090	0.0234	0.3838	2.6058
	Endokrines Gewebe	0.0358	0.0301	1.1887	0.8413
10	Gastrointestinal	0.0019	0.0000	undef	0.0000
	Gehirn	0.0007	0.0010	0.7200	1.3890
	Haematopoetisch	0.0000	0.0000	undef	undef
	Haut	0.0037	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0065	0.0000	undef
15	Herz	0.0244	0.0000	undef	0.0000
	Hoden	0.0000	0.0000	undef	undef
	Lunge	0.0374	0.0491	0.7621	1.3122
	Magen-Speiserohre	0.0290	0.0077	3.7816	0.2644
	Muskel-Skelett	0.0000	0.0000	undef	undef
20	Niere	0.0027	0.0068	0.3965	2.5219
	Pankreas	0.0017	0.2209	0.0075	133.7133
	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0065	0.0085	0.7677	1.3026
	Uterus Endometrium	0.0000	0.0000	undef	undef
25	Uterus Myometrium	0.0152	0.0000	undef	0.0000
	Uterus allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0192			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0267			
30	Sinnesorgane	0.0000			
	Weisse Blutkoerperchen	0.0000			
	Zervix	0.0106			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0000
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0667
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.1224
	Eierstock_n	0.0000
55	Eierstock_t	0.1013
	Endokrines Gewebe	0.0000
	Foetal	0.0268
	Gastrointestinal	0.0122
	Haematopoetisch	0.0057
60	Haut-Muskel	0.0065
	Hoden	0.0000
	Lunge	0.0246
	Nerven	0.0000
	Prostata	0.0205
65	Sinnesorgane	0.0000
	Uterus_n	0.0125

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haefufigkeit	%Haefufigkeit	N/T	T/N	
Blase	0.0039	0.0537	0.0726	13.7665	5
Brust	0.0077	0.0207	0.3712	2.6940	
Duenn darm	0.0368	0.0000	undef	0.0000	
Eierstock	0.0150	0.0078	1.9188	0.5212	
Endokrines Gewebe	0.0102	0.0100	1.0189	0.9815	
Gastrointestinal	0.0421	0.0093	4.5559	0.2195	10
Gehirn	0.0118	0.0195	0.6063	1.6494	
Haematopoetisch	0.0174	0.0379	0.4587	2.1798	
Haut	0.0110	0.0000	undef	0.0000	
Hepatisch	0.0048	0.0518	0.0919	10.8799	
Herz	0.0127	0.0275	0.4626	2.1618	15
Hoden	0.0115	0.0117	0.9839	1.0163	
Lunge	0.0114	0.0061	1.8628	0.5368	
Magen-Speiserohre	0.0000	0.0460	0.0000	undef	
Muskel-Skelett	0.0154	0.0060	2.5700	0.3891	
Niere	0.0054	0.0068	0.7930	1.2610	20
Pankreas	0.0050	0.0331	0.1496	6.6857	
Penis	0.0090	0.0533	0.1685	5.9357	
Prostata	0.0174	0.0192	0.9099	1.0990	
Uterus_Endometrium	0.0068	0.0000	undef	0.0000	25
Uterus_Myometrium	0.0152	0.0000	undef	0.0000	
Uterus_allgemein	0.0204	0.0000	undef	0.0000	
Brust-Hyperplasie	0.0064				
Prostata-Hyperplasie	0.0238				
Samenblase	0.0000				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0251				30
Zervix	0.0106				

	FOETUS	
	%Haefufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0167	
Gehirn	0.0438	
Haematopoetisch	0.0118	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0107	
Lunge	0.0181	
Nebenniere	0.0000	
Niere	0.0247	45
Placenta	0.0061	
Prostata	0.0249	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haefufigkeit	
Brust	0.0408	50
Eierstock_n	0.0000	
Eierstock_t	0.0101	
Endokrines_Gewebe	0.0000	55
Foetal	0.0087	
Gastrointestinal	0.0122	
Haematopoetisch	0.0000	
Haut-Muskel	0.0130	60
Hoden	0.0000	
Lunge	0.0082	
Nerven	0.0010	
Prostata	0.0068	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0195	0.2556	0.0763	13.1109
	Brust	0.0166	0.0357	0.4656	2.1477
	Duenn darm	0.0061	0.0662	0.0927	10.7893
	Eierstock	0.0389	0.0052	7.4832	0.1336
	Endokrines Gewebe	0.0392	0.0326	1.2017	0.8321
10	Gastrointestinal	0.0019	0.0000	undef	0.0000
	Gehirn	0.0007	0.0606	0.0122	81.9491
	Haematopoetisch	0.0107	0.0000	undef	0.0000
	Haut	0.0220	0.5085	0.0433	23.0839
	Hepatisch	0.0238	0.0518	0.4596	2.1760
15	Herz	0.0085	0.0000	undef	0.0000
	Hoden	0.0115	0.0000	undef	0.0000
	Lunge	0.0104	0.0041	2.5402	0.3937
	Magen-Speiserohre	0.0000	0.0077	0.0000	undef
	Muskel-Skelett	0.0600	0.0480	1.2493	0.8005
20	Niere	0.0407	0.0068	5.9478	0.1681
	Pankreas	0.0198	0.0331	0.5983	1.6714
	Penis	0.0030	0.1066	0.0281	35.6140
	Prostata	0.0000	0.0021	0.0000	undef
	Uterus_Endometrium	0.0405	0.0000	undef	0.0000
	Uterus_Myometrium	0.0305	0.1155	0.2641	3.7870
25	Uterus_allgemein	0.0153	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0064			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0000			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0000			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0696
	Gastrointestinal	0.3332
	Gehirn	0.0000
	Haematopoetisch	0.2202
40	Haut	0.0000
	Hepatisch	1.6381
	Herz-Blutgefasse	0.0285
	Lunge	0.1337
	Nebenniere	1.0903
45	Niere	0.6301
	Placenta	0.6786
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0544
	Eierstock_n	0.0000
55	Eierstock_t	0.1063
	Endokrines_Gewebe	0.0000
	Foetal	0.4264
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0030
	Prostata	0.0000
65	Sinnesorgane	0.0000
	Uterus_n	0.0250

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0195	0.2301	0.0847 11.7998	5
Brust	0.0192	0.0113	1.7013 0.5878	
Duennndarm	0.0061	0.0331	0.1854 5.3946	
Eierstock	0.0180	0.0000	undef 0.0000	
Endokrines_Gewebe	0.0034	0.1555	0.0219 45.6387	
Gastrointestinal	0.0000	0.0370	0.0000 undef	10
Gehirn	0.0214	0.1561	0.1374 7.2801	
Haematopoetisch	0.0134	0.0000	undef 0.0000	
Haut	0.0073	0.0000	undef 0.0000	
Hepatisch	0.0048	0.0000	undef 0.0000	
Herz	0.0085	0.0000	undef 0.0000	
Hoden	0.0000	0.0000	undef undef	15
Lunge	0.0197	0.0020	9.6527 0.1036	
Magen-Speiserohre	0.0000	0.0077	0.0000 undef	
Muskel-Skelett	0.0034	0.1320	0.0260 38.5221	
Niere	0.0109	0.0068	1.5861 0.6305	
Pankreas	0.0083	0.0000	undef 0.0000	20
Penis	0.0000	0.0000	undef undef	
Prostata	0.0000	0.0021	0.0000 undef	
Uterus_Endometrium	0.0203	0.0000	undef 0.0000	
Uterus_Myometrium	0.0000	0.0475	0.0000 undef	
Uterus_allgemein	0.0153	0.0000	undef 0.0000	25
Brust-Hyperplasie	0.0128			
Prostata-Hyperplasie	0.0030			
Samenblase	0.0000			
Sinnesorgane	0.0235			
Weisse_Blutkoerperchen	0.0000			30
Zervix	0.0213			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0417	35
Gastrointestinal	0.1361	
Gehirn	0.0063	
Haematopoetisch	0.1337	
Haut	0.0000	40
Hepatisch	0.3380	
Herz-Blutgefuesse	0.0249	
Lunge	0.0578	
Nebenniere	0.5071	
Niere	0.2594	45
Placenta	0.4120	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0612	
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0000	55
Foetal	0.1188	
Gastrointestinal	0.0244	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0154	60
Lunge	0.0000	
Nerven	0.0141	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0416	65

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase 0.0039	0.0383	0.1017	9.8332
	Brust 0.0077	0.0075	1.0208	0.9796
	Duenndarm 0.0061	0.0496	0.1236	8.0920
	Eierstock 0.0000	0.0130	0.0000	undef
	Endokrines Gewebe 0.0153	0.0000	undef	0.0000
10	Gastrointestinal 0.0115	0.0000	undef	0.0000
	Gehirn 0.0022	0.0031	0.7200	1.3890
	Haematopoetisch 0.0013	0.0000	undef	0.0000
	Haut 0.0073	0.0000	undef	0.0000
	Hepatisch 0.0048	0.0129	0.3676	2.7200
15	Herz 0.0000	0.0000	undef	undef
	Hoden 0.0000	0.0000	undef	undef
	Lunge 0.0042	0.0102	0.4064	2.4605
	Magen-Speiserohre 0.0290	0.0077	3.7816	0.2644
	Muskel-Skelett 0.0000	0.0000	undef	undef
20	Niere 0.0136	0.0000	undef	0.0000
	Pankreas 0.0017	0.0000	undef	0.0000
	Penis 0.0030	0.0000	undef	0.0000
	Prostata 0.0065	0.0106	0.6142	1.6282
	Uterus_Endometrium 0.0068	0.0528	0.1280	7.8106
25	Uterus_Myometrium 0.0000	0.0000	undef	undef
	Uterus_allgemein 0.0000	0.0000	undef	undef
	Brust-Hyperplasie 0.0032			
	Prostata-Hyperplasie 0.0268			
	Samenblase 0.0267			
	Sinnesorgane 0.0235			
30	Weisse_Blutkoerperchen 0.0000			
	Zervix 0.0319			

	FOETUS
	%Haeufigkeit
35	Entwicklung 0.0139
	Gastrointestinal 0.0139
	Gehirn 0.0000
	Haematopoetisch 0.0039
40	Haut 0.0000
	Hepatisch 0.0000
	Herz-Blutgefuesse 0.0000
	Lunge 0.0036
	Nebenniere 0.0254
45	Niere 0.0062
	Placenta 0.0000
	Prostata 0.0000
	Sinnesorgane 0.0000

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
	%Haeufigkeit
50	Brust 0.0204
	Eierstock_n 0.0000
55	Eierstock_t 0.0203
	Endokrines_Gewebe 0.0245
	Foetal 0.0128
	Gastrointestinal 0.0122
	Haematopoetisch 0.0000
60	Haut-Muskel 0.0000
	Hoden 0.0154
	Lunge 0.0082
	Nerven 0.0090
	Prostata 0.0068
65	Sinnesorgane 0.0000
	Uterus_n 0.0000

	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0039	0.0358	0.1090	9.1777	5
Brust	0.0077	0.0207	0.3712	2.6940	
Duendarm	0.0215	0.0331	0.6488	1.5413	
Eierstock	0.0030	0.0078	0.3838	2.6058	
Endokrines_Gewebe	0.0000	0.0000	undef	undef	
Gastrointestinal	0.0575	0.0046	12.4251	0.0805	10
Gehirn	0.0000	0.0072	0.0000	undef	
Haematopoetisch	0.0214	0.0000	undef	0.0000	
Haut	0.0037	0.0000	undef	0.0000	
Hepatisch	0.0000	0.0194	0.0000	undef	
Herz	0.0000	0.0137	0.0000	undef	
Hoden	0.0000	0.0585	0.0000	undef	15
Lunge	0.0104	0.0225	0.4618	2.1652	
Magen-Speiserohre	0.0000	0.0000	undef	undef	
Muskel-Skelett	0.0257	0.0120	2.1416	0.4669	
Niere	0.0054	0.0000	undef	0.0000	
Pankreas	0.0099	0.0221	0.4487	2.2286	20
Penis	0.0000	0.0000	undef	undef	
Prostata	0.0022	0.0000	undef	0.0000	
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0000	undef	undef	
Uterus_allgemein	0.0000	0.0000	undef	undef	25
Brust-Hyperplasie	0.0096				
Prostata-Hyperplasie	0.0000				
Samenblase	0.0000				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0000				30
Zervix	0.0000				

FOETUS					
	%Haeufigkeit				
Entwicklung	0.0000				35
Gastrointestinal	0.0000				
Gehirn	0.0000				
Haematopoetisch	0.0000				
Haut	0.0000				40
Hepatisch	0.0000				
Herz-Blutgefuesse	0.0000				
Lunge	0.0000				
Nebenniere	0.0000				
Niere	0.0000				45
Placenta	0.0000				
Prostata	0.0000				
Sinnesorgane	0.0000				

NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN					
	%Haeufigkeit				
Brust	0.0136				50
Eierstock_n	0.0000				
Eierstock_t	0.0051				
Endokrines_Gewebe	0.0000				55
Foetal	0.0000				
Gastrointestinal	0.0122				
Haematopoetisch	0.0000				
Haut-Muskel	0.0000				
Hoden	0.0000				60
Lunge	0.0164				
Nerven	0.0000				
Prostata	0.0000				
Sinnesorgane	0.0000				
Uterus_n	0.0000				65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0351	0.3144	0.1116	8.9591
	Brust	0.0281	0.0470	0.5989	1.6698
	Duendarm	0.0092	0.0662	0.1390	7.1929
	Eierstock	0.0569	0.0208	2.7342	0.3657
	Endokrines_Gewebe	0.0596	0.0527	1.1321	0.8833
10	Gastrointestinal	0.0019	0.0139	0.1381	7.2434
	Gehirn	0.0022	0.0945	0.0235	42.5950
	Haematopoetisch	0.0174	0.0000	undef	0.0000
	Haut	0.0220	0.5085	0.0433	23.0839
	Hepatisch	0.0285	0.0582	0.4902	2.0400
15	Herz	0.0223	0.0000	undef	0.0000
	Hoden	0.0173	0.0000	undef	0.0000
	Lunge	0.0145	0.0082	1.7781	0.5624
	Magen-Speiserohre	0.0290	0.0077	3.7816	0.2644
	Muskel-Skelett	0.0788	0.0540	1.4595	0.6852
20	Niere	0.0489	0.0137	3.5687	0.2802
	Pankreas	0.0264	0.0442	0.5983	1.6714
	Penis	0.0090	0.1066	0.0842	11.8713
	Prostata	0.0000	0.0064	0.0000	undef
	Uterus_Endometrium	0.1013	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0381	0.1494	0.2551	3.9206
	Uterus_allgemein	0.0153	0.0954	0.1601	6.2452
	Brust-Hyperplasie	0.0096			
	Prostata-Hyperplasie	0.0059			
	Samenblase	0.0000			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0000			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0696
	Gastrointestinal	0.4554
	Gehirn	0.0000
	Haematopoetisch	0.2753
40	Haut	0.0000
	Hepatisch	1.6381
	Herz-Blutgefuesse	0.0605
	Lunge	0.1879
	Nebenniere	1.1663
45	Niere	0.8215
	Placenta	0.7816
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0612
	Eierstock_n	0.0000
55	Eierstock_t	0.1164
	Endokrines_Gewebe	0.0000
	Foetal	0.4665
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0030
	Prostata	0.0068
65	Sinnesorgane	0.0000
	Uterus_n	0.0291

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0039	0.0332	0.1173 8.5221	5
Brust	0.0038	0.0263	0.1458 6.8574	
Duenndarm	0.0184	0.0331	0.5561 1.7982	
Eierstock	0.0150	0.0156	0.9594 1.0423	
Endokrines Gewebe	0.0170	0.0075	2.2642 0.4417	
Gastrointestinal	0.0192	0.0324	0.5917 1.6901	10
Gehirn	0.0133	0.0113	1.1781 0.8488	
Haematopoetisch	0.0174	0.0000	undef 0.0000	
Haut	0.0220	0.0000	undef 0.0000	
Hepatisch	0.0048	0.0388	0.1225 8.1599	
Herz	0.0339	0.0137	2.4671 0.4053	
Hoden	0.0288	0.0234	1.2299 0.8130	15
Lunge	0.0218	0.0184	1.1854 0.8436	
Magen-Speiserohre	0.0000	0.0307	0.0000 undef	
Muskel-Skelett	0.0291	0.0240	1.2136 0.8240	
Niere	0.0136	0.0137	0.9913 1.0088	
Pankreas	0.0050	0.0166	0.2991 3.3428	20
Penis	0.0120	0.0000	undef 0.0000	
Prostata	0.0153	0.0149	1.0236 0.9769	
Uterus_Endometrium	0.0203	0.0000	undef 0.0000	
Uterus_Myometrium	0.0000	0.0204	0.0000 undef	
Uterus_allgemein	0.0051	0.0000	undef 0.0000	25
Brust-Hyperplasie	0.0128			
Prostata-Hyperplasie	0.0178			
Samenblase	0.0267			
Sinnesorgane	0.0118			
Weisse_Blutkoerperchen	0.0243			30
Zervix	0.0213			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0139	35
Gastrointestinal	0.0056	
Gehirn	0.0125	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0320	
Lunge	0.0036	
Nebenniere	0.0254	
Niere	0.0124	45
Placenta	0.0121	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0204	50
Eierstock_n	0.0000	
Eierstock_t	0.0354	
Endokrines_Gewebe	0.0000	55
Foetal	0.0105	
Gastrointestinal	0.0000	
Haematopoetisch	0.0171	
Haut-Muskel	0.0454	
Hoden	0.0000	60
Lunge	0.0246	
Nerven	0.0211	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0195	0.1457	0.1338	7.4732
	Brust	0.0013	0.0169	0.0756	13.2250
	Duenn darm	0.0061	0.0496	0.1236	8.0920
	Eierstock	0.0090	0.0260	0.3454	2.8954
	Endokrines Gewebe	0.0119	0.0125	0.9509	1.0516
10	Gastrointestinal	0.1648	0.0231	7.1237	0.1404
	Gehirn	0.0030	0.0308	0.0960	10.4173
	Haematopoetisch	0.2099	0.0000	undef	0.0000
	Haut	0.0000	0.0000	undef	undef
	Hepatisch	0.0000	0.2006	0.0000	undef
15	Herz	0.0032	0.0137	0.2313	4.3235
	Hoden	0.0000	0.0819	0.0000	undef
	Lunge	0.0956	0.2965	0.3223	3.1023
	Magen-Speiserohre	0.0097	0.1917	0.0504	19.8329
	Muskel-Skelett	0.0959	0.0240	3.9977	0.2501
20	Niere	0.0081	0.0205	0.3965	2.5219
	Pankreas	0.0182	0.0497	0.3656	2.7350
	Penis	0.0030	0.1333	0.0225	44.5175
	Prostata	0.0131	0.0064	2.0473	0.4885
	Uterus_Endometrium	0.0000	0.0000	undef	undef
25	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0051	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0096			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0267			
	Sinnesorgane	0.0706			
30	Weisse_Blutkoerperchen	0.0017			
	Zervix	0.0000			

35	FOETUS	
	%Haeufigkeit	
	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0000
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

50	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
	Brust	0.0068
	Eierstock_n	0.0000
55	Eierstock_t	0.2076
	Endokrines_Gewebe	0.0000
	Foetal	0.0006
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0082
	Nerven	0.0030
	Prostata	0.0068
65	Sinnesorgane	0.0000
	Uterus_n	0.0042

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0039	0.0281	0.1387 7.2110	5
Brust	0.0217	0.0132	1.6527 0.6051	
Duennndarm	0.0123	0.0000	undef 0.0000	
Eierstock	0.0000	0.0026	0.0000 undef	
Endokrines_Gewebe	0.0102	0.0150	0.6792 1.4722	
Gastrointestinal	0.0115	0.0046	2.4850 0.4024	10
Gehirn	0.0148	0.0154	0.9599 1.0417	
Haematopoetisch	0.0107	0.0000	undef 0.0000	
Haut	0.0220	0.0847	0.2599 3.8473	
Hepatisch	0.0000	0.0000	undef undef	
Herz	0.0085	0.0000	undef 0.0000	
Hoden	0.0058	0.0117	0.4920 2.0326	15
Lunge	0.0145	0.0061	2.3708 0.4218	
Magen-Speiserohre	0.0000	0.0077	0.0000 undef	
Muskel-Skelett	0.0086	0.0360	0.2380 4.2024	
Niere	0.0081	0.0000	undef 0.0000	
Pankreas	0.0050	0.0166	0.2991 3.3428	20
Penis	0.0030	0.0000	undef 0.0000	
Prostata	0.0109	0.0064	1.7060 0.5862	
Uterus_Endometrium	0.0203	0.0000	undef 0.0000	
Uterus_Myometrium	0.0076	0.0068	1.1223 0.8911	25
Uterus_allgemein	0.0102	0.0000	undef 0.0000	
Brust-Hyperplasie	0.0096			
Prostata-Hyperplasie	0.0059			
Samenblase	0.0089			
Sinnesorgane	0.0000			
Weisse_Blutkoerperchen	0.0286			30
Zervix	0.0106			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0028	
Gehirn	0.0125	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0036	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0061	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0204	
Eierstock_n	0.0000	
Eierstock_t	0.0051	
Endokrines_Gewebe	0.0000	55
Foetal	0.0029	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0000	
Nerven	0.0010	
Prostata	0.0068	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0039	0.0256	0.1525	6.5555
	Brust	0.0038	0.0056	0.6805	1.4694
	Duenn darm	0.0031	0.0000	undef	0.0000
	Eierstock	0.0210	0.0052	4.0294	0.2482
	Endokrines Gewebe	0.0017	0.0000	undef	0.0000
10	Gastrointestinal	0.0057	0.0000	undef	0.0000
	Gehirn	0.0037	0.0031	1.1999	0.8334
	Haematopoetisch	0.0040	0.0000	undef	0.0000
	Haut	0.0073	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0000	undef	undef
15	Herz	0.0000	0.0000	undef	undef
	Hoden	0.0000	0.0000	undef	undef
	Lunge	0.0073	0.0020	3.5562	0.2812
	Magen-Speiserohre	0.0000	0.0000	undef	undef
	Muskel-Skelett	0.0069	0.0000	undef	0.0000
20	Niere	0.0027	0.0000	undef	0.0000
	Pankreas	0.0017	0.0000	undef	0.0000
	Penis	0.0030	0.0000	undef	0.0000
	Prostata	0.0000	0.0021	0.0000	undef
	Uterus_Endometrium	0.0000	0.0000	undef	undef
	Uterus_Myometrium	0.0076	0.0068	1.1223	0.8911
25	Uterus_allgemein	0.0102	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0089			
	Sinnesorgane	0.0000			
30	Weisse_Blutkoerperchen	0.0121			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0139
	Gehirn	0.0063
	Haematopoetisch	0.0039
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0036
	Lunge	0.0036
	Nebenniere	0.0000
45	Niere	0.0062
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.0000
55	Eierstock_t	0.0101
	Endokrines_Gewebe	0.0000
	Foetal	0.0029
	Gastrointestinal	0.0488
	Haematopoetisch	0.0114
60	Haut-Muskel	0.0130
	Hoden	0.0154
	Lunge	0.0000
	Nerven	0.0020
	Prostata	0.0274
65	Sinnesorgane	0.0155
	Uterus_n	0.0083

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0039	0.0256	0.1525 6.5555	5
Brust	0.0115	0.0150	0.7656 1.3062	
Duenn darm	0.0307	0.0000	undef 0.0000	
Eierstock	0.0150	0.0052	2.8781 0.3474	
Endokrines Gewebe	0.0119	0.0150	0.7925 1.2619	
Gastrointestinal	0.0057	0.0139	0.4142 2.4145	10
Gehirn	0.0074	0.0092	0.8000 1.2501	
Haematopoetisch	0.0147	0.0000	undef 0.0000	
Haut	0.0037	0.1695	0.0217 46.1678	
Hepatisch	0.0048	0.0000	undef 0.0000	
Herz	0.0074	0.0137	0.5397 1.8529	15
Hoden	0.0000	0.0234	0.0000 undef	
Lunge	0.0104	0.0082	1.2701 0.7873	
Magen-Speiserohre	0.0000	0.0230	0.0000 undef	
Muskel-Skelett	0.0086	0.0000	undef 0.0000	
Niere	0.0190	0.0000	undef 0.0000	20
Pankreas	0.0066	0.0055	1.1966 0.8357	
Penis	0.0150	0.0267	0.5616 1.7807	
Prostata	0.0087	0.0149	0.5849 1.7096	
Uterus_Endometrium	0.0203	0.0000	undef 0.0000	
Uterus_Myometrium	0.0000	0.0136	0.0000 undef	25
Uterus_allgemein	0.0051	0.0000	undef 0.0000	
Brust-Hyperplasie	0.0064			
Prostata-Hyperplasie	0.0178			
Samenblase	0.0178			
Sinnesorgane	0.0235			
Weisse_Blutkoerperchen	0.0104			30
Zervix	0.0000			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0111	
Gehirn	0.0125	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0260	
Herz-Blutgefuesse	0.0071	
Lunge	0.0217	
Nebenniere	0.0254	
Niere	0.0062	45
Placenta	0.0061	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0136	
Eierstock_n	0.0000	
Eierstock_t	0.0101	
Endokrines_Gewebe	0.0245	55
Foetal	0.0181	
Gastrointestinal	0.0244	
Haematopoetisch	0.0114	
Haut-Muskel	0.0065	
Hoden	0.0000	60
Lunge	0.0164	
Nerven	0.0090	
Prostata	0.0068	
Sinnesorgane	0.0232	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0312	0.1968	0.1585	6.3096
	Brust	0.0281	0.0526	0.5347	1.8702
	Duenn darm	0.2177	0.0000	undef	0.0000
	Eierstock	0.0000	0.0494	0.0000	undef
	Endokrines Gewebe	0.0017	0.0000	undef	0.0000
10	Gastrointestinal	0.2778	0.0324	8.5792	0.1166
	Gehirn	0.0007	0.0133	0.0554	18.0566
	Haematopoetisch	0.1470	0.0000	undef	0.0000
	Haut	0.0551	0.0000	undef	0.0000
	Hepatisch	0.0000	0.1876	0.0000	undef
15	Herz	0.0011	0.0137	0.0771	12.9706
	Hoden	0.0000	0.2339	0.0000	undef
	Lunge	0.0623	0.1206	0.5166	1.9356
	Magen-Speiserohre	0.0000	0.0153	0.0000	undef
	Muskel-Skelett	0.1216	0.0300	4.0548	0.2466
20	Niere	0.0081	0.1027	0.0793	12.6097
	Pankreas	0.0743	0.2209	0.3365	2.9714
	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0065	0.0149	0.4387	2.2795
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
	Uterus_Myometrium	0.0000	0.0000	undef	undef
25	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0384			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0000			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0017			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0068
	Eierstock_n	0.0000
	Eierstock_t	0.0051
55	Endokrines Gewebe	0.0000
	Foetal	0.0000
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0082
	Nerven	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000
65	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0273	0.1713	0.1594 6.2745	5
Brust	0.0217	0.0395	0.5509 1.8152	
Duenndarm	0.1901	0.0000	undef 0.0000	
Eierstock	0.0000	0.0390	0.0000 undef	
Endokrines_Gewebe	0.0017	0.0000	undef 0.0000	
Gastrointestinal	0.2491	0.0278	8.9737 0.1114	10
Gehirn	0.0000	0.0113	0.0000 undef	
Haematopoetisch	0.1297	0.0000	undef 0.0000	
Haut	0.0477	0.0000	undef 0.0000	
Hepatisch	0.0000	0.1423	0.0000 undef	
Herz	0.0032	0.0000	undef 0.0000	
Hoden	0.0000	0.2105	0.0000 undef	15
Lunge	0.0457	0.0818	0.5588 1.7894	
Magen-Speiserohre	0.0000	0.0153	0.0000 undef	
Muskel-Skelett	0.0822	0.0240	3.4266 0.2918	
Niere	0.0054	0.1027	0.0529 18.9146	20
Pankreas	0.0694	0.2430	0.2855 3.5020	
Penis	0.0000	0.0000	undef undef	
Prostata	0.0153	0.0106	1.4331 0.6978	
Uterus_Endometrium	0.0068	0.0000	undef 0.0000	
Uterus_Myometrium	0.0000	0.0000	undef undef	
Uterus_allgemein	0.0000	0.0000	undef undef	25
Brust-Hyperplasie	0.0288			
Prostata-Hyperplasie	0.0030			
Samenblase	0.0000			
Sinnesorgane	0.0118			
Weisse_Blutkoerperchen	0.0009			30
Zervix	0.0000			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0000	
Haematopoetisch	0.0079	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0000	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0068	
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0000	55
Foetal	0.0000	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0082	
Nerven	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0858	0.4525	0.1896	5.2742
	Brust	0.1279	0.0977	1.3087	0.7641
	Duennndarm	0.4998	0.2150	2.3242	0.4302
	Eierstock	0.0180	0.1353	0.1328	7.5280
	Endokrines_Gewebe	0.0307	0.0176	1.7466	0.5725
10	Gastrointestinal	0.7434	0.1341	5.5413	0.1805
	Gehirn	0.0030	0.0524	0.0565	17.7093
	Haematopoetisch	0.6884	0.0000	undef	0.0000
	Haut	0.0551	0.1695	0.3249	3.0779
	Hepatisch	0.0381	0.7635	0.0499	20.0598
15	Herz	0.0095	0.0412	0.2313	4.3235
	Hoden	0.0000	0.1403	0.0000	undef
	Lunge	0.2773	0.5418	0.5119	1.9536
	Magen-Speiserohre	0.0676	0.3450	0.1961	5.0999
	Muskel-Skelett	0.3203	0.0660	4.8544	0.2060
20	Niere	0.0163	0.1780	0.0915	10.9284
	Pankreas	0.0529	0.2319	0.2279	4.3875
	Penis	0.0000	0.0000	undef	undef
	Prostata	0.0828	0.0192	4.3220	0.2314
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
	Uterus_Myometrium	0.0076	0.0204	0.3741	2.6732
25	Uterus_allgemein	0.0102	0.0000	undef	0.0000
	Brust-Hyperplasie	0.1343			
	Prostata-Hyperplasie	0.0268			
	Samenblase	0.0267			
	Sinnesorgane	0.0353			
30	Weisse_Blutkoerperchen	0.0043			
	Zervix	0.0852			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0118
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.1837
	Eierstock_n	0.0000
	Eierstock_t	0.0405
55	Endokrines_Gewebe	0.0000
	Foetal	0.0023
	Gastrointestinal	0.0976
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0020
	Prostata	0.0000
65	Sinnesorgane	0.0000
	Uterus_n	0.0042

	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0039	0.0204	0.1907	5.2444	5
Brust	0.0077	0.0056	1.3611	0.7347	
Duendarm	0.0184	0.0331	0.5561	1.7982	
Eierstock	0.0150	0.0104	1.4391	0.6949	
Endokrines_Gewebe	0.0102	0.0000	undef	0.0000	
Gastrointestinal	0.0019	0.0046	0.4142	2.4145	10
Gehirn	0.0074	0.0072	1.0285	0.9723	
Haematopoetisch	0.0067	0.0000	undef	0.0000	
Haut	0.0147	0.0000	undef	0.0000	
Hepatisch	0.0000	0.0065	0.0000	undef	
Herz	0.0064	0.0000	undef	0.0000	
Hoden	0.0345	0.0234	1.4759	0.6775	15
Lunge	0.0073	0.0143	0.5080	1.9684	
Magen-Speiserohre	0.0000	0.0000	undef	undef	
Muskel-Skelett	0.0034	0.0000	undef	0.0000	
Niere	0.0054	0.0000	undef	0.0000	
Pankreas	0.0050	0.0055	0.8974	1.1143	20
Penis	0.0030	0.0000	undef	0.0000	
Prostata	0.0153	0.0043	3.5827	0.2791	
Uterus_Endometrium	0.0068	0.0000	undef	0.0000	
Uterus_Myometrium	0.0000	0.0068	0.0000	undef	
Uterus_allgemein	0.0000	0.0000	undef	undef	25
Brust-Hyperplasie	0.0000				
Prostata-Hyperplasie	0.0089				
Samenblase	0.0178				
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0069				30
Zervix	0.0106				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0111	
Gehirn	0.0000	
Haematopoetisch	0.0039	
Haut	0.2513	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0071	
Lunge	0.0036	
Nebenniere	0.0000	
Niere	0.0062	45
Placenta	0.0061	
Prostata	0.0249	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0136	
Eierstock_n	0.0000	
Eierstock_t	0.0152	
Endokrines_Gewebe	0.0000	55
Foetal	0.0041	
Gastrointestinal	0.0122	
Haematopoetisch	0.0000	
Haut-Muskel	0.0130	
Hoden	0.0077	60
Lunge	0.0000	
Nerven	0.0040	
Prostata	0.0137	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0039	0.0204	0.1907	5.2444
	Brust	0.0141	0.0282	0.4991	2.0038
	Duenn darm	0.0061	0.0000	undef	0.0000
	Eierstock	0.0090	0.0312	0.2878	3.4745
	Endokrines Gewebe	0.0153	0.0176	0.8733	1.1451
10	Gastrointestinal	0.0192	0.0231	0.8283	1.2072
	Gehirn	0.0222	0.0113	1.9635	0.5093
	Haematopoetisch	0.0094	0.0000	undef	0.0000
	Haut	0.0110	0.0000	undef	0.0000
	Hepatisch	0.0048	0.0129	0.3676	2.7200
15	Herz	0.0159	0.0000	undef	0.0000
	Hoden	0.0000	0.0234	0.0000	undef
	Lunge	0.0135	0.0123	1.1007	0.9085
	Magen-Speiserohre	0.0097	0.0153	0.6303	1.5866
	Muskel-Skelett	0.0188	0.0300	0.6282	1.5918
20	Niere	0.0163	0.0205	0.7930	1.2610
	Pankreas	0.0083	0.0221	0.3739	2.6743
	Penis	0.0090	0.0267	0.3369	2.9678
	Prostata	0.0109	0.0043	2.5591	0.3908
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
	Uterus_Myometrium	0.0076	0.0204	0.3741	2.6732
25	Uterus_allgemein	0.0102	0.0954	0.1067	9.3678
	Brust-Hyperplasie	0.0032			
	Prostata-Hyperplasie	0.0119			
	Samenblase	0.0089			
	Sinnesorgane	0.0235			
30	Weisse_Blutkoerperchen	0.0087			
	Zervix	0.0000			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0111
	Gehirn	0.0000
	Haematopoetisch	0.0039
40	Haut	0.0000
	Hepatisch	0.0260
	Herz-Blutgefuesse	0.0107
	Lunge	0.0072
	Nebenniere	0.0000
45	Niere	0.0124
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.1595
	Eierstock_t	0.0203
55	Endokrines_Gewebe	0.0000
	Foetal	0.0058
	Gastrointestinal	0.0488
	Haematopoetisch	0.0114
60	Haut-Muskel	0.0032
	Hoden	0.0154
	Lunge	0.0164
	Nerven	0.0060
	Prostata	0.0068
	Sinnesorgane	0.0000
65	Uterus_n	0.0083

	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0429	0.2173	0.1974	5.0656	5
Brust	0.0409	0.0357	1.1462	0.8725	
Duennndarm	0.1870	0.0662	2.8269	0.3537	
Eierstock	0.0030	0.0833	0.0360	27.7957	
Endokrines_Gewebe	0.0034	0.0050	0.6792	1.4722	
Gastrointestinal	0.2740	0.0740	3.7016	0.2702	10
Gehirn	0.0007	0.0216	0.0343	29.1683	
Haematopoetisch	0.2165	0.0000	undef	0.0000	
Haut	0.0220	0.0000	undef	0.0000	
Hepatisch	0.0095	0.3429	0.0277	36.0397	
Herz	0.0074	0.0275	0.2698	3.7059	
Hoden	0.0000	0.0585	0.0000	undef	15
Lunge	0.1039	0.1738	0.5977	1.6731	
Magen-Speiserohre	0.0387	0.0920	0.4202	2.3799	
Muskel-Skelett	0.0994	0.0420	2.3660	0.4227	
Niere	0.0054	0.0753	0.0721	13.8707	
Pankreas	0.0231	0.1160	0.1994	5.0142	20
Penis	0.0000	0.0000	undef	undef	
Prostata	0.0262	0.0000	undef	0.0000	
Uterus_Endometrium	0.0068	0.0000	undef	0.0000	
Uterus_Myometrium	0.0076	0.0000	undef	0.0000	
Uterus_allgemein	0.0051	0.0000	undef	0.0000	25
Brust-Hyperplasie	0.0512				
Prostata-Hyperplasie	0.0178				
Samenblase	0.0089				
Sinnesorgane	0.0118				
Weisse_Blutkoerperchen	0.0000				30
Zervix	0.0319				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0000	
Haematopoetisch	0.0079	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0000	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0612	
Eierstock_n	0.0000	
Eierstock_t	0.0152	
Endokrines_Gewebe	0.0000	55
Foetal	0.0012	
Gastrointestinal	0.0366	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0000	
Nerven	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0117	0.0588	0.1990	5.0259
	Brust	0.0102	0.0094	1.0888	0.9184
	Duenn darm	0.0153	0.0331	0.4634	2.1579
	Eierstock	0.0539	0.0130	4.1445	0.2413
	Endokrines Gewebe	0.0187	0.0075	2.4906	0.4015
10	Gastrointestinal	0.0134	0.0093	1.4496	0.6898
	Gehirn	0.0133	0.0123	1.0799	0.9260
	Haematopoetisch	0.0134	0.0000	undef	0.0000
	Haut	0.0147	0.0000	undef	0.0000
	Hepatisch	0.0000	0.0129	0.0000	undef
15	Herz	0.0170	0.0137	1.2336	0.8107
	Hoden	0.0173	0.0234	0.7380	1.3551
	Lunge	0.0062	0.0020	3.0482	0.3281
	Magen-Speiserohre	0.0676	0.0000	undef	0.0000
	Muskel-Skelett	0.0171	0.0180	0.9518	1.0506
20	Niere	0.0109	0.0274	0.3965	2.5219
	Pankreas	0.0066	0.0110	0.5983	1.6714
	Penis	0.0269	0.0533	0.5054	1.9786
	Prostata	0.0327	0.0213	1.5354	0.6513
	Uterus Endometrium	0.0068	0.0000	undef	0.0000
	Uterus Myometrium	0.0534	0.0000	undef	0.0000
25	Uterus allgemein	0.0306	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0000			
	Sinnesorgane	0.0118			
30	Weisse Blutkoerperchen	0.0061			
	Zervix	0.0213			

		FOETUS
		%Haeufigkeit.
35	Entwicklung	0.0139
	Gastrointestinal	0.0555
	Gehirn	0.0000
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0260
	Herz-Blutgefuesse	0.0071
	Lunge	0.0253
	Nebenniere	0.0254
45	Niere	0.0000
	Placenta	0.0485
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0204
	Eierstock_n	0.0000
	Eierstock_t	0.2430
55	Endokrines Gewebe	0.0245
	Foetal	0.0338
	Gastrointestinal	0.0122
	Haematopoetisch	0.0171
60	Haut-Muskel	0.0680
	Hoden	0.0077
	Lunge	0.0000
	Nerven	0.0151
	Prostata	0.0342
	Sinnesorgane	0.0000
65	Uterus_n	0.1166

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0273	0.1227	0.2225 4.4952	5
Brust	0.0243	0.0263	0.9236 1.0828	
Duennndarm	0.1625	0.0000	undef 0.0000	
Eierstock	0.0000	0.0364	0.0000 undef	
Endokrines_Gewebe	0.0000	0.0000	undef undef	
Gastrointestinal	0.2146	0.0093	23.1935 0.0431	10
Gehirn	0.0000	0.0062	0.0000 undef	
Haematopoetisch	0.1203	0.0000	undef 0.0000	
Haut	0.0441	0.0000	undef 0.0000	
Hepatisch	0.0000	0.1229	0.0000 undef	
Herz	0.0021	0.0000	undef 0.0000	
Hoden	0.0000	0.0935	0.0000 undef	15
Lunge	0.0353	0.0900	0.3926 2.5473	
Magen-Speiserohre	0.0000	0.0077	0.0000 undef	
Muskel-Skelett	0.0857	0.0120	7.1388 0.1401	
Niere	0.0081	0.0753	0.1081 9.2471	
Pankreas	0.0529	0.1381	0.3829 2.6116	20
Penis	0.0000	0.0000	undef undef	
Prostata	0.0109	0.0064	1.7060 0.5862	
Uterus_Endometrium	0.0000	0.0000	undef undef	
Uterus_Myometrium	0.0000	0.0000	undef undef	
Uterus_allgemein	0.0000	0.0000	undef undef	25
Brust-Hyperplasie	0.0160			
Prostata-Hyperplasie	0.0000			
Samenblase	0.0000			
Sinnesorgane	0.0118			
Weisse_Blutkoerperchen	0.0009			30
Zervix	0.0106			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0000	
Haematopoetisch	0.0039	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0000	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0000	
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0000	55
Foetal	0.0000	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0082	
Nerven	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0078	0.0332	0.2347	4.2611
	Brust	0.0153	0.0169	0.9074	1.1021
	Duennndarm	0.0092	0.0165	0.5561	1.7982
	Eierstock	0.0210	0.0130	1.6118	0.6204
	Endokrines_Gewebe	0.0085	0.0150	0.5660	1.7667
10	Gastrointestinal	0.0153	0.0231	0.6627	1.5090
	Gehirn	0.0185	0.0226	0.8181	1.2223
	Haematopoetisch	0.0227	0.0379	0.5999	1.6669
	Haut	0.0073	0.0000	undef	0.0000
	Hepatisch	0.0048	0.0194	0.2451	4.0800
15	Herz	0.0201	0.0137	1.4649	0.6827
	Hoden	0.0115	0.0234	0.4920	2.0326
	Lunge	0.0166	0.0164	1.0161	0.9842
	Magen-Speiserohre	0.0000	0.0153	0.0000	undef
	Muskel-Skelett	0.0137	0.0060	2.2844	0.4378
20	Niere	0.0081	0.0274	0.2974	3.3626
	Pankreas	0.0050	0.0055	0.8974	1.1143
	Penis	0.0180	0.0267	0.6739	1.4839
	Prostata	0.0065	0.0106	0.6142	1.6282
	Uterus_Endometrium	0.0135	0.0528	0.2561	3.9053
	Uterus_Myometrium	0.0152	0.0408	0.3741	2.6732
25	Uterus_allgemein	0.0458	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0256			
	Prostata-Hyperplasie	0.0208			
	Samenblase	0.0178			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0061			
	Zervix	0.0106			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0417
	Gastrointestinal	0.0167
	Gehirn	0.0250
	Haematopoetisch	0.0079
40	Haut	0.2513
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0142
	Lunge	0.0181
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0242
	Prostata	0.0249
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0136
	Eierstock_n	0.0000
	Eierstock_t	0.0101
55	Endokrines_Gewebe	0.0000
	Foetal	0.0082
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0292
	Hoden	0.0077
	Lunge	0.0164
	Nerven	0.0110
	Prostata	0.0068
	Sinnesorgane	0.0000
65	Uterus_n	0.0208

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0078	0.0332	0.2347	4.2611	5
Brust	0.0026	0.0000	undef	0.0000	
Duenn darm	0.0000	0.0000	undef	undef	
Eierstock	0.0000	0.0000	undef	undef	10
Endokrines Gewebe	0.0000	0.0000	undef	undef	
Gastrointestinal	0.0115	0.0000	undef	0.0000	
Gehirn	0.0000	0.0000	undef	undef	15
Haematopoetisch	0.0147	0.0000	undef	0.0000	
Haut	0.0000	0.0000	undef	undef	
Hepatisch	0.0000	0.0000	undef	undef	20
Herz	0.0000	0.0000	undef	undef	
Hoden	0.0000	0.0000	undef	undef	
Lunge	0.0021	0.0020	1.0161	0.9842	25
Magen-Speiserohre	0.0000	0.0000	undef	undef	
Muskel-Skelett	0.0069	0.0000	undef	0.0000	
Niere	0.0000	0.0000	undef	undef	30
Pankreas	0.0017	0.0607	0.0272	36.7712	
Penis	0.0000	0.0000	undef	undef	
Prostata	0.0000	0.0000	undef	undef	35
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0000	undef	undef	
Uterus_allgemein	0.0000	0.0000	undef	undef	40
Brust-Hyperplasie	0.0000				
Prostata-Hyperplasie	0.0000				
Samenblase	0.0000				45
Sinnesorgane	0.0000				
Weisse_Blutkoerperchen	0.0009				
Zervix	0.0000				50

FOETUS				
	%Haeufigkeit			
Entwicklung	0.0000			35
Gastrointestinal	0.0000			
Gehirn	0.0000			
Haematopoetisch	0.0039			
Haut	0.0000			40
Hepatisch	0.0000			
Herz-Blutgefuesse	0.0000			
Lunge	0.0000			
Nebenniere	0.0000			
Niere	0.0000			45
Placenta	0.0000			
Prostata	0.0000			
Sinnesorgane	0.0000			

NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN				
	%Haeufigkeit			
Brust	0.0000			50
Eierstock_n	0.0000			
Eierstock_t	0.0000			
Endokrines_Gewebe	0.0000			55
Foetal	0.0000			
Gastrointestinal	0.0000			
Haematopoetisch	0.0000			
Haut-Muskel	0.0000			
Hoden	0.0000			60
Lunge	0.0000			
Nerven	0.0000			
Prostata	0.0000			
Sinnesorgane	0.0000			
Uterus_n	0.0000			65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0624	0.2607	0.2393	4.1791
	Brust	0.0051	0.0282	0.1815	5.5104
	Duennndarm	0.0337	0.0000	undef	0.0000
	Eierstock	0.0030	0.0937	0.0320	31.2702
	Endokrines Gewebe	0.0034	0.0025	1.3585	0.7361
10	Gastrointestinal	0.1360	0.0694	1.9604	0.5101
	Gehirn	0.0015	0.0318	0.0464	21.5290
	Haematopoetisch	0.1337	0.0000	undef	0.0000
	Haut	0.0147	0.0000	undef	0.0000
	Hepatisch	0.0000	0.4594	0.0000	undef
15	Herz	0.0127	0.0137	0.9252	1.0809
	Hoden	0.0000	0.1754	0.0000	undef
	Lunge	0.0540	0.2086	0.2590	3.8610
	Magen-Speiserohre	0.0193	0.0767	0.2521	3.9666
	Muskel-Skelett	0.1747	0.0240	7.2815	0.1373
20	Niere	0.0190	0.0205	0.9252	1.0808
	Pankreas	0.0611	0.0773	0.7906	1.2649
	Penis	0.0030	0.0000	undef	0.0000
	Prostata	0.0196	0.0021	9.2126	0.1085
	Uterus Endometrium	0.0135	0.0000	undef	0.0000
	Uterus Myometrium	0.0000	0.0000	undef	undef
25	Uterus allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0064			
	Prostata-Hyperplasie	0.0089			
	Samenblase	0.0089			
	Sinnesorgane	0.0353			
30	Weisse Blutkoerperchen	0.0009			
	Zervix	0.0426			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0000
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0061
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0000
	Eierstock_n	0.0000
	Eierstock_t	0.0101
55	Endokrines Gewebe	0.0000
	Foetal	0.0000
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0010
	Prostata	0.0000
	Sinnesorgane	0.0000
65	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0702	0.2761	0.2542	3.9333	5
Brust	0.0614	0.0620	0.9899	1.0102	
Duennndarm	0.2024	0.1489	1.3594	0.7356	
Eierstock	0.0060	0.0442	0.1354	7.3832	
Endokrines_Gewebe	0.0255	0.0025	10.1887	0.0981	
Gastrointestinal	0.4138	0.0740	5.5913	0.1788	10
Gehirn	0.0044	0.0431	0.1029	9.7228	
Haematopoetisch	0.5774	0.0000	undef	0.0000	
Haut	0.0257	0.0000	undef	0.0000	
Hepatisch	0.0285	0.3364	0.0848	11.7866	
Herz	0.0095	0.0412	0.2313	4.3235	15
Hoden	0.0000	0.2689	0.0000	undef	
Lunge	0.1735	0.2678	0.6476	1.5441	
Magen-Speiseroehre	0.0870	0.1917	0.4538	2.2037	
Muskel-Skelett	0.2193	0.0180	12.1835	0.0821	
Niere	0.0136	0.1164	0.1166	8.5746	
Pankreas	0.0677	0.1822	0.3717	2.6906	20
Penis	0.0030	0.1333	0.0225	44.5175	
Prostata	0.0327	0.0085	3.8386	0.2605	
Uterus_Endometrium	0.0000	0.0000	undef	undef	
Uterus_Myometrium	0.0000	0.0000	undef	undef	
Uterus_allgemein	0.0051	0.0000	undef	0.0000	25
Brust-Hyperplasie	0.0927				
Prostata-Hyperplasie	0.0178				
Samenblase	0.0267				
Sinnesorgane	0.2823				
Weisse_Blutkoerperchen	0.0087				30
Zervix	0.0426				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0056	
Gehirn	0.0000	
Haematopoetisch	0.0157	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0000	
Nebenniere	0.0254	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.4082	
Eierstock_n	0.0000	
Eierstock_t	0.3493	
Endokrines_Gewebe	0.0000	55
Foetal	0.0082	
Gastrointestinal	0.1953	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0491	
Nerven	0.0030	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0195	0.0690	0.2825	3.5400
	Brust	0.0166	0.0320	0.5204	1.9216
	Duennndarm	0.0031	0.0000	undef	0.0000
	Eierstock	0.0150	0.0130	1.1513	0.8686
	Endokrines_Gewebe	0.0085	0.0100	0.8491	1.1778
10	Gastrointestinal	0.0019	0.0093	0.2071	4.8289
	Gehirn	0.0067	0.0442	0.1507	6.6362
	Haematopoetisch	0.0187	0.0000	undef	0.0000
	Haut	0.0073	0.0000	undef	0.0000
	Hepatisch	0.0238	0.0194	1.2255	0.8160
15	Herz	0.0625	0.1512	0.4135	2.4182
	Hoden	0.0345	0.0117	2.9518	0.3388
	Lunge	0.0322	0.0286	1.1249	0.8889
	Magen-Speiserohre	0.0000	0.0307	0.0000	undef
	Muskel-Skelett	0.0668	0.1260	0.5303	1.8857
	Niere	0.0190	0.0342	0.5551	1.8014
20	Pankreas	0.0050	0.1160	0.0427	23.3998
	Penis	0.0299	0.0000	undef	0.0000
	Prostata	0.0131	0.0170	0.7677	1.3026
	Uterus_Endometrium	0.0068	0.0528	0.1280	7.8106
	Uterus_Myometrium	0.0305	0.0204	1.4964	0.6683
25	Uterus_allgemein	0.0357	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0000			
	Prostata-Hyperplasie	0.0208			
	Samenblase	0.0356			
	Sinnesorgane	0.0588			
30	Weisse_Blutkoerperchen	0.0000			
	Zervix	0.0319			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0000
	Gastrointestinal	0.0167
	Gehirn	0.0000
	Haematopoetisch	0.0118
40	Haut	0.0000
	Hepatisch	0.0520
	Herz-Blutgefuesse	0.0107
	Lunge	0.0325
	Nebenniere	0.0000
45	Niere	0.0494
	Placenta	0.0909
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0340
	Eierstock_n	0.0000
	Eierstock_t	0.0203
55	Endokrines_Gewebe	0.0490
	Foetal	0.0297
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
	Haut-Muskel	0.0000
60	Hoden	0.0154
	Lunge	0.0082
	Nerven	0.0030
	Prostata	0.0000
	Sinnesorgane	0.0000
65	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0390	0.1355	0.2878 3.4744	5
Brust	0.0269	0.0564	0.4764 2.0992	
Duennndarm	0.1932	0.0000	undef 0.0000	
Eierstock	0.0000	0.0312	0.0000 undef	
Endokrines_Gewebe	0.0017	0.0025	0.6792 1.4722	
Gastrointestinal	0.2203	0.0509	4.3300 0.2309	10
Gehirn	0.0000	0.0103	0.0000 undef	
Haematopoetisch	0.0882	0.0000	undef 0.0000	
Haut	0.0184	0.0000	undef 0.0000	
Hepatisch	0.0000	0.0518	0.0000 undef	
Herz	0.0095	0.0137	0.6939 1.4412	15
Hoden	0.0000	0.0702	0.0000 undef	
Lunge	0.0478	0.1472	0.3246 3.0809	
Magen-Speiserohre	0.0193	0.0153	1.2605 0.7933	
Muskel-Skelett	0.1011	0.0120	8.4237 0.1187	
Niere	0.0054	0.0548	0.0991 10.0878	20
Pankreas	0.0099	0.0552	0.1795 5.5714	
Penis	0.0000	0.2933	0.0000 undef	
Prostata	0.0153	0.0021	7.1654 0.1396	
Uterus_Endometrium	0.0068	0.0000	undef 0.0000	
Uterus_Myometrium	0.0000	0.0068	0.0000 undef	
Uterus_allgemein	0.0000	0.0000	undef undef	25
Brust-Hyperplasie	0.0192			
Prostata-Hyperplasie	0.0119			
Samenblase	0.0000			
Sinnesorgane	0.0000			
Weisse_Blutkoerperchen	0.0009			30
Zervix	0.0213			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0000	
Gehirn	0.0000	
Haematopoetisch	0.0079	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0000	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0000	50
Eierstock_n	0.0000	
Eierstock_t	0.0000	
Endokrines_Gewebe	0.0000	55
Foetal	0.0000	
Gastrointestinal	0.0000	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0000	60
Lunge	0.0000	
Nerven	0.0000	
Prostata	0.0000	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0273	0.0895	0.3051	3.2777
	Brust	0.0665	0.0808	0.8230	1.2151
	Duennndarm	0.0429	0.0000	undef	0.0000
	Eierstock	0.0539	0.0234	2.3025	0.4343
	Endokrines_Gewebe	0.0630	0.0978	0.6444	1.5518
10	Gastrointestinal	0.0441	0.0324	1.3608	0.7348
	Gehirn	0.0554	0.1006	0.5510	1.8149
	Haematopoetisch	0.0454	0.0379	1.1998	0.8335
	Haut	0.0257	0.0000	undef	0.0000
	Hepatisch	0.0381	0.0453	0.8403	1.1900
15	Herz	0.0435	0.0825	0.5268	1.8981
	Hoden	0.0575	0.0000	undef	0.0000
	Lunge	0.1008	0.0552	1.8252	0.5479
	Magen-Speiserohre	0.0580	0.0997	0.5818	1.7188
	Muskel-Skelett	0.0976	0.0660	1.4797	0.6758
20	Niere	0.0516	0.0890	0.5795	1.7255
	Pankreas	0.0248	0.0773	0.3205	3.1200
	Penis	0.0599	0.1066	0.5616	1.7807
	Prostata	0.0567	0.0766	0.7393	1.3527
	Uterus_Endometrium	0.0405	0.1055	0.3841	2.6035
	Uterus_Myometrium	0.0534	0.0475	1.1223	0.8911
25	Uterus_allgemein	0.0866	0.0000	undef	0.0000
	Brust-Hyperplasie	0.0416			
	Prostata-Hyperplasie	0.0654			
	Samenblase	0.0712			
	Sinnesorgane	0.0823			
30	Weisse_Blutkoerperchen	0.1110			
	Zervix	0.0319			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0139
	Gastrointestinal	0.0361
	Gehirn	0.0125
	Haematopoetisch	0.0433
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0071
	Lunge	0.0253
	Nebenniere	0.0254
45	Niere	0.0432
	Placenta	0.0364
	Prostata	0.0499
	Sinnesorgane	0.0126

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.1020
	Eierstock_n	0.1595
	Eierstock_t	0.0709
55	Endokrines_Gewebe	0.0000
	Foetal	0.0635
	Gastrointestinal	0.0000
	Haematopoetisch	0.0114
60	Haut-Muskel	0.0680
	Hoden	0.0463
	Lunge	0.0328
	Nerven	0.0351
	Prostata	0.0342
	Sinnesorgane	0.0464
65	Uterus_n	0.0083

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0195	0.0639	0.3051 3.2777	5
Brust	0.0345	0.0470	0.7350 1.3606	
Duennndarm	0.0399	0.1985	0.2008 4.9797	
Eierstock	0.0150	0.0676	0.2214 4.5168	
Endokrines_Gewebe	0.0238	0.0878	0.2717 3.6805	
Gastrointestinal	0.0862	0.1064	0.8103 1.2341	10
Gehirn	0.0067	0.0277	0.2400 4.1669	
Haematopoetisch	0.0080	0.0000	undef 0.0000	
Haut	0.0587	0.0000	undef 0.0000	
Hepatisch	0.0190	0.0323	0.5882 1.7000	
Herz	0.0011	0.0962	0.0110 90.7941	15
Hoden	0.0000	0.0234	0.0000 undef	
Lunge	0.0062	0.0164	0.3810 2.6245	
Magen-Speiseroehre	0.0387	0.3450	0.1120 8.9248	
Muskel-Skelett	0.0000	0.0360	0.0000 undef	
Niere	0.0760	0.1643	0.4626 2.1617	20
Pankreas	0.0677	0.0276	2.4530 0.4077	
Penis	0.0090	0.0533	0.1685 5.9357	
Prostata	0.0109	0.0255	0.4265 2.3446	
Uterus_Endometrium	0.0270	0.0000	undef 0.0000	
Uterus_Myometrium	0.0076	0.0272	0.2806 3.5642	25
Uterus_allgemein	0.0000	0.4771	0.0000 undef	
Brust-Hyperplasie	0.0576			
Prostata-Hyperplasie	0.0119			
Samenblase	0.1068			
Sinnesorgane	0.0235			
Weisse_Blutkoerperchen	0.0061			30
Zervix	0.0319			

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0111	
Gehirn	0.0813	
Haematopoetisch	0.0000	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0000	
Lunge	0.0145	
Nebenniere	0.0000	
Niere	0.0309	45
Placenta	0.0121	
Prostata	0.0249	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0136	
Eierstock_n	0.0000	
Eierstock_t	0.0101	
Endokrines_Gewebe	0.0000	55
Foetal	0.0122	
Gastrointestinal	0.4149	
Haematopoetisch	0.0000	
Haut-Muskel	0.0000	
Hoden	0.0154	60
Lunge	0.0573	
Nerven	0.0040	
Prostata	0.0068	
Sinnesorgane	0.0000	
Uterus_n	0.0000	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0156	0.0511	0.3051	3.2777
	Brust	0.0166	0.0320	0.5204	1.9216
	Duenn darm	0.0828	0.0000	undef	0.0000
	Eierstock	0.0030	0.0416	0.0720	13.8979
	Endokrines_Gewebe	0.0017	0.0050	0.3396	2.9444
10	Gastrointestinal	0.1226	0.0278	4.4178	0.2264
	Gehirn	0.0000	0.0123	0.0000	undef
	Haematopoetisch	0.0481	0.0000	undef	0.0000
	Haut	0.0294	0.0000	undef	0.0000
	Hepatisch	0.0000	0.1294	0.0000	undef
15	Herz	0.0032	0.0137	0.2313	4.3235
	Hoden	0.0000	0.1403	0.0000	undef
	Lunge	0.0447	0.1390	0.3213	3.1128
	Magen-Speiserohre	0.0097	0.0000	undef	0.0000
	Muskel-Skelett	0.0497	0.0060	8.2810	0.1208
20	Niere	0.0054	0.0685	0.0793	12.6097
	Pankreas	0.0215	0.0166	1.2963	0.7714
	Penis	0.0030	0.0000	undef	0.0000
	Prostata	0.0044	0.0043	1.0236	0.9769
	Uterus_Endometrium	0.0000	0.0000	undef	undef
25	Uterus_Myometrium	0.0000	0.0000	undef	undef
	Uterus_allgemein	0.0000	0.0000	undef	undef
	Brust-Hyperplasie	0.0352			
	Prostata-Hyperplasie	0.0030			
	Samenblase	0.0000			
	Sinnesorgane	0.0000			
30	Weisse_Blutkoerperchen	0.0000			
	Zervix	0.0106			

		FOETUS
		%Haeufigkeit.
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0079
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0000
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0136
	Eierstock_n	0.0000
	Eierstock_t	0.0000
55	Endokrines_Gewebe	0.0000
	Foetal	0.0000
	Gastrointestinal	0.0000
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0000
	Nerven	0.0000
	Prostata	0.0000
65	Sinnesorgane	0.0000
	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0117	0.0383	0.3051	3.2777	5
Brust	0.0179	0.0207	0.8661	1.1546	
Duenndarm	0.0245	0.0165	1.4830	0.6743	
Eierstock	0.0329	0.0156	2.1106	0.4738	
Endokrines_Gewebe	0.0221	0.0326	0.6792	1.4722	
Gastrointestinal	0.0153	0.0139	1.1045	0.9054	10
Gehirn	0.0296	0.0288	1.0285	0.9723	
Haematopoetisch	0.0187	0.0379	0.4940	2.0241	
Haut	0.0257	0.0000	undef	0.0000	
Hepatisch	0.0048	0.0129	0.3676	2.7200	
Herz	0.0201	0.0550	0.3662	2.7306	15
Hoden	0.0058	0.0117	0.4920	2.0326	
Lunge	0.0218	0.0225	0.9699	1.0311	
Magen-Speiserohre	0.0387	0.0383	1.0084	0.9916	
Muskel-Skelett	0.0171	0.0240	0.7139	1.4008	
Niere	0.0190	0.0068	2.7756	0.3603	20
Pankreas	0.0116	0.0221	0.5235	1.9102	
Penis	0.0090	0.0000	undef	0.0000	
Prostata	0.0327	0.0319	1.0236	0.9769	
Uterus_Endometrium	0.0068	0.0000	undef	0.0000	
Uterus_Myometrium	0.0000	0.0136	0.0000	undef	25
Uterus_allgemein	0.0560	0.0000	undef	0.0000	
Brust-Hyperplasie	0.0032				
Prostata-Hyperplasie	0.0208				
Samenblase	0.0178				
Sinnesorgane	0.0118				
Weisse_Blutkoerperchen	0.0260				30
Zervix	0.0000				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.0000	35
Gastrointestinal	0.0139	
Gehirn	0.0313	
Haematopoetisch	0.0039	
Haut	0.0000	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0071	
Lunge	0.0217	
Nebenniere	0.0000	
Niere	0.0000	45
Placenta	0.0424	
Prostata	0.0249	
Sinnesorgane	0.0000	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0408	50
Eierstock_n	0.0000	
Eierstock_t	0.0405	
Endokrines_Gewebe	0.0245	55
Foetal	0.0087	
Gastrointestinal	0.0000	
Haematopoetisch	0.0057	
Haut-Muskel	0.0097	
Hoden	0.0231	60
Lunge	0.0082	
Nerven	0.0251	
Prostata	0.0205	
Sinnesorgane	0.0000	
Uterus_n	0.0125	65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.1170	0.3067	0.3814	2.6222
	Brust	0.3019	0.2387	1.2646	0.7908
	Duennndarm	1.1559	0.0000	undef	0.0000
	Eierstock	0.0120	0.0676	0.1771	5.6460
	Endokrines Gewebe	0.0034	0.0125	0.2717	3.6805
10	Gastrointestinal	1.2798	0.1804	7.0940	0.1410
	Gehirn	0.0007	0.0380	0.0195	51.3918
	Haematopoetisch	0.4785	0.0000	undef	0.0000
	Haut	0.1322	0.0000	undef	0.0000
	Hepatisch	0.0285	0.4594	0.0621	16.0932
15	Herz	0.0138	0.0275	0.5011	1.9955
	Hoden	0.0000	0.2456	0.0000	undef
	Lunge	0.3625	0.3435	1.0554	0.9475
	Magen-Speiserohre	0.0483	0.1533	0.3151	3.1733
	Muskel-Skelett	0.2124	0.0480	4.4260	0.2259
20	Niere	0.0163	0.1917	0.0850	11.7691
	Pankreas	0.1074	0.4528	0.2371	4.2171
	Penis	0.0000	0.0533	0.0000	undef
	Prostata	0.0806	0.0255	3.1562	0.3168
	Uterus_Endometrium	0.0068	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0152	0.0000	undef	0.0000
	Uterus_allgemein	0.0102	0.0000	undef	0.0000
	Brust-Hyperplasie	0.3805			
	Prostata-Hyperplasie	0.0505			
	Samenblase	0.0356			
	Sinnesorgane	0.5175			
30	Weisse_Blutkoerperchen	0.0078			
	Zervix	0.0213			

		FOETUS
		%Haeufigkeit.
35	Entwicklung	0.0000
	Gastrointestinal	0.0000
	Gehirn	0.0000
	Haematopoetisch	0.0039
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0000
	Lunge	0.0000
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.0303
	Prostata	0.0000
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.7687
	Eierstock_n	0.0000
	Eierstock_t	0.1924
55	Endokrines Gewebe	0.0000
	Foetal	0.0029
	Gastrointestinal	0.0976
	Haematopoetisch	0.0000
60	Haut-Muskel	0.0000
	Hoden	0.0000
	Lunge	0.0328
	Nerven	0.0020
	Prostata	0.0000
	Sinnesorgane	0.0000
65	Uterus_n	0.0000

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	NORMAL	TUMOR	Verhaeltnisse		
	%Haeufigkeit	%Haeufigkeit	N/T	T/N	
Blase	0.0195	0.0486	0.4014	2.4911	5
Brust	0.0550	0.1297	0.4241	2.3580	
Duennndarm	0.0123	0.1654	0.0741	13.4866	
Eierstock	0.0449	0.0728	0.6167	1.6214	
Endokrines_Gewebe	0.0119	0.0075	1.5849	0.6309	
Gastrointestinal	0.0153	0.1018	0.1506	6.6398	10
Gehirn	0.0037	0.0359	0.1029	9.7228	
Haematopoetisch	0.0134	0.0000	undef	0.0000	
Haut	0.0404	0.0847	0.4765	2.0985	
Hepatisch	0.0285	0.0323	0.8824	1.1333	
Herz	0.0233	0.0825	0.2827	3.5374	15
Hoden	0.0460	0.0351	1.3119	0.7622	
Lunge	0.0551	0.0491	1.1219	0.8913	
Magen-Speiserohre	0.0000	0.0613	0.0000	undef	
Muskel-Skelett	0.1148	0.3120	0.3679	2.7180	
Niere	0.0271	0.0137	1.9826	0.5044	20
Pankreas	0.0116	0.0884	0.1309	7.6408	
Penis	0.0180	0.0533	0.3369	2.9678	
Prostata	0.0087	0.0319	0.2730	3.6634	
Uterus_Endometrium	0.0270	0.0000	undef	0.0000	
Uterus_Myometrium	0.0610	0.1902	0.3206	3.1187	25
Uterus_allgemein	0.0509	0.0000	undef	0.0000	
Brust-Hyperplasie	0.0288				
Prostata-Hyperplasie	0.0089				
Samenblase	0.0178				
Sinnesorgane	0.0118				
Weisse_Blutkoerperchen	0.0000				30
Zervix	0.0532				

	FOETUS	
	%Haeufigkeit	
Entwicklung	0.1809	35
Gastrointestinal	0.1388	
Gehirn	0.0063	
Haematopoetisch	0.0551	
Haut	0.2513	40
Hepatisch	0.0000	
Herz-Blutgefuesse	0.0605	
Lunge	0.1409	
Nebenniere	0.2535	
Niere	0.1235	45
Placenta	0.0364	
Prostata	0.0748	
Sinnesorgane	1.3934	

	NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN	
	%Haeufigkeit	
Brust	0.0204	
Eierstock_n	0.0000	
Eierstock_t	0.0304	
Endokrines_Gewebe	0.0000	55
Foetal	0.0315	
Gastrointestinal	0.0122	
Haematopoetisch	0.0000	
Haut-Muskel	0.0518	
Hoden	0.0231	60
Lunge	0.0000	
Nerven	0.0040	
Prostata	0.0137	
Sinnesorgane	0.0000	
Uterus_n	0.0083	65

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		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0624	0.1380	0.4520	2.2125
	Brust	0.0345	0.0564	0.6125	1.6327
	Duennndarm	0.0491	0.0165	2.9659	0.3372
	Eierstock	0.0689	0.0520	1.3239	0.7553
	Endokrines_Gewebe	0.0392	0.0276	1.4202	0.7041
10	Gastrointestinal	0.0460	0.0648	0.7100	1.4084
	Gehirn	0.0333	0.0678	0.4909	2.0372
	Haematopoetisch	0.0374	0.0758	0.4940	2.0241
	Haut	0.0257	0.1695	0.1516	6.5954
	Hepatisch	0.1142	0.0518	2.2059	0.4533
15	Herz	0.0774	0.7010	0.1104	9.0616
	Hoden	0.2589	0.1520	1.7030	0.5872
	Lunge	0.0540	0.0491	1.1007	0.9085
	Magen-Speiserohre	0.1256	0.2070	0.6069	1.6477
	Muskel-Skelett	0.1542	0.2100	0.7343	1.3619
20	Niere	0.0109	0.0959	0.1133	8.8268
	Pankreas	0.0198	0.1270	0.1561	6.4071
	Penis	0.0359	0.0800	0.4493	2.2259
	Prostata	0.0785	0.0511	1.5354	0.6513
	Uterus_Endometrium	0.0338	0.0528	0.6402	1.5621
25	Uterus_Myometrium	0.0457	0.0679	0.6734	1.4851
	Uterus_allgemein	0.0764	0.0954	0.8006	1.2490
	Brust-Hyperplasie	0.0224			
	Prostata-Hyperplasie	0.0624			
	Samenblase	0.0445			
	Sinnesorgane	0.0118			
30	Weisse_Blutkoerperchen	0.0529			
	Zervix	0.0532			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0139
	Gastrointestinal	0.0167
	Gehirn	0.0000
	Haematopoetisch	0.0039
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefasse	0.0249
	Lunge	0.0108
	Nebenniere	0.0000
45	Niere	0.0000
	Placenta	0.3333
	Prostata	0.1995
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.1156
	Eierstock_n	0.0000
	Eierstock_t	0.1873
55	Endokrines_Gewebe	0.0000
	Foetal	0.0181
	Gastrointestinal	0.0000
	Haematopoetisch	0.0057
60	Haut-Muskel	0.0486
	Hoden	0.0000
	Lunge	0.0328
	Nerven	0.0020
	Prostata	0.0274
65	Sinnesorgane	0.0155
	Uterus_n	0.0541

	NORMAL	TUMOR	Verhaeltnisse	
	%Haeufigkeit	%Haeufigkeit	N/T T/N	
Blase	0.0585	0.1278	0.4576 2.1852	5
Brust	0.0742	0.0507	1.4619 0.6841	
Duennndarm	0.1134	0.0000	undef 0.0000	
Eierstock	0.0659	0.0390	1.6885 0.5922	
Endokrines_Gewebe	0.0324	0.0176	1.8437 0.5424	
Gastrointestinal	0.0690	0.0694	0.9940 1.0060	10
Gehirn	0.0325	0.0893	0.3641 2.7464	
Haematopoetisch	0.0842	0.0000	undef 0.0000	
Haut	0.0404	0.0000	undef 0.0000	
Hepatisch	0.1855	0.0647	2.8677 0.3487	
Herz	0.0435	0.0962	0.4516 2.2145	15
Hoden	0.0230	0.0585	0.3936 2.5408	
Lunge	0.2275	0.1063	2.1396 0.4674	
Magen-Speiserohre	0.0483	0.1993	0.2424 4.1252	
Muskel-Skelett	0.0857	0.0720	1.1898 0.8405	
Niere	0.0706	0.0274	2.5774 0.3880	20
Pankreas	0.0347	0.2927	0.1185 8.4367	
Penis	0.0090	0.0267	0.3369 2.9678	
Prostata	0.0458	0.0319	1.4331 0.6978	
Uterus_Endometrium	0.0270	0.0000	undef 0.0000	
Uterus_Myometrium	0.0610	0.0204	2.9927 0.3341	25
Uterus_allgemein	0.0917	0.0000	undef 0.0000	
Brust-Hyperplasie	0.0416			
Prostata-Hyperplasie	0.0386			
Samenblase	0.0267			
Sinnesorgane	0.1647			
Weisse_Blutkoerperchen	0.2497			30
Zervix	0.0000			
FOETUS				
	%Haeufigkeit			35
Entwicklung	0.0000			
Gastrointestinal	0.0916			
Gehirn	0.0500			
Haematopoetisch	0.0865			
Haut	0.0000			40
Hepatisch	0.0520			
Herz-Blutgefuesse	0.0178			
Lunge	0.0434			
Nebenniere	0.0000			
Niere	0.0432			45
Placenta	0.1636			
Prostata	0.0000			
Sinnesorgane	0.0000			
NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN				
	%Haeufigkeit			50
Brust	0.1156			
Eierstock_n	0.0000			
Eierstock_t	0.0354			
Endokrines_Gewebe	0.0490			55
Foetal	0.0571			
Gastrointestinal	0.0854			
Haematopoetisch	0.0057			
Haut-Muskel	0.0097			
Hoden	0.0154			60
Lunge	0.0491			
Nerven	0.0562			
Prostata	0.0615			
Sinnesorgane	0.0000			
Uterus_n	0.1957			65

		NORMAL	TUMOR	Verhaeltnisse	
		%Haeufigkeit	%Haeufigkeit	N/T	T/N
5	Blase	0.0546	0.1099	0.4967	2.0135
	Brust	0.0563	0.0489	1.1517	0.8683
	Duenndarm	0.1380	0.0331	4.1708	0.2398
	Eierstock	0.0599	0.0650	0.9210	1.0858
	Endokrines_Gewebe	0.0238	0.0351	0.6792	1.4722
10	Gastrointestinal	0.1322	0.1804	0.7328	1.3647
	Gehirn	0.0229	0.0452	0.5072	1.9714
	Haematopoetisch	0.0241	0.0000	undef	0.0000
	Haut	0.1689	0.0000	undef	0.0000
	Hepatisch	0.0285	0.0518	0.5515	1.8133
15	Herz	0.1166	0.1649	0.7067	1.4150
	Hoden	0.0115	0.0117	0.9839	1.0163
	Lunge	0.1070	0.1329	0.8050	1.2422
	Magen-Speiserohre	0.1450	0.0613	2.3635	0.4231
	Muskel-Skelett	0.0685	0.0240	2.8555	0.3502
20	Niere	0.0570	0.0753	0.7570	1.3210
	Pankreas	0.0165	0.1491	0.1108	9.0256
	Penis	0.0779	0.0267	2.9202	0.3424
	Prostata	0.0610	0.0255	2.3885	0.4187
	Uterus_Endometrium	0.0338	0.0000	undef	0.0000
25	Uterus_Myometrium	0.0991	0.0340	2.9179	0.3427
	Uterus_allgemein	0.0509	0.1908	0.2669	3.7471
	Brust-Hyperplasie	0.0064			
	Prostata-Hyperplasie	0.0386			
	Samenblase	0.0801			
	Sinnesorgane	0.0588			
30	Weisse_Blutkoerperchen	0.0616			
	Zervix	0.1810			

		FOETUS
		%Haeufigkeit
35	Entwicklung	0.0139
	Gastrointestinal	0.0194
	Gehirn	0.0000
	Haematopoetisch	0.0275
40	Haut	0.0000
	Hepatisch	0.0000
	Herz-Blutgefuesse	0.0142
	Lunge	0.0145
	Nebenniere	0.0000
45	Niere	0.0247
	Placenta	0.0364
	Prostata	0.0499
	Sinnesorgane	0.0000

		NORMIERTE/SUBTRAHIERTE BIBLIOTHEKEN
		%Haeufigkeit
50	Brust	0.0068
	Eierstock_n	0.0000
	Eierstock_t	0.0101
55	Endokrines_Gewebe	0.0000
	Foetal	0.0064
	Gastrointestinal	0.0976
	Haematopoetisch	0.0057
60	Haut-Muskel	0.0259
	Hoden	0.0309
	Lunge	0.1802
	Nerven	0.0050
	Prostata	0.0274
	Sinnesorgane	0.0000
65	Uterus_n	0.0125

2.2 Fisher-Test

Um zu entscheiden, ob eine Partial-Sequenz S eines Gens in einer Bibliothek für Normal-Gewebe signifikant häufiger oder seltener vorkommt als in einer Bibliothek für entartetes Gewebe, wird Fishers Exakter Test, ein statistisches Standardverfahren (Hays, W. L., (1991) Statistics, Harcourt Brace College Publishers, Fort Worth), durchgeführt.

Die Null-Hypothese lautet: die beiden Bibliotheken können bezüglich der Häufigkeit zu S homologer Sequenzen nicht unterschieden werden. Falls die Null-Hypothese mit hinreichend hoher Sicherheit abgelehnt werden kann, wird das zu S gehörende Gen als interessanter Kandidat für ein Krebs-Gen akzeptiert, und es wird im nächsten Schritt versucht, eine Verlängerung seiner Sequenz zu erreichen.

Beispiel 3

Automatische Verlängerung der Partial-Sequenz

Die automatische Verlängerung der Partial-Sequenz S vollzieht sich in drei Schritten:

1. Ermittlung aller zu S homologen Sequenzen aus der Gesamtmenge der zur Verfügung stehenden Sequenzen mit Hilfe von BLAST
2. Assemblierung dieser Sequenzen mittels des Standardprogramms GAP4 (Bonfield, J. K., Smith, K. F., und Staden R. (1995), Nucleic Acids Research 23 4992-4999) (Contig-Bildung).
3. Berechnung einer Konsens-Sequenz C aus den assemblierten Sequenzen.

Die Konsens-Sequenz C wird im allgemeinen länger sein als die Ausgangssequenz s. Ihr elektronischer Northern-Blot wird demzufolge von dem für S abweichen. Ein erneuter Fisher-Test entscheidet, ob die Alternativ-Hypothese der Abweichung von einer gleichmäßigen Expression in beiden Bibliotheken aufrechterhalten werden kann. Ist dies der Fall, wird versucht, C in gleicher Weise wie S zu verlängern. Diese Iteration wird mit der jeweils erhaltenen Konsensus-Sequenzen C_i (i: Index der Iteration) fortgesetzt, bis die Alternativ-Hypothese verworfen wird (if H_0 Exit; Abbruchkriterium I) oder bis keine automatische Verlängerung mehr möglich ist (while $C_i > C_{i-1}$; Abbruchkriterium II).

Im Fall des Abbruchkriteriums II bekommt man mit der nach der letzten Iteration vorliegenden Konsens-Sequenz eine komplette oder annähernd komplette Sequenz eines Gens, das mit hoher statistischer Sicherheit mit Krebs in Zusammenhang gebracht werden kann.

Analog der oben beschriebenen Beispiele konnten die in der Tabelle I beschriebenen Nukleinsäure-Sequenzen aus Blasentumorgewebe gefunden werden.

Ferner konnten zu den einzelnen Nukleinsäure-Sequenzen die Peptidsequenzen (ORF's) bestimmt werden, die in der Tabelle II aufgelistet sind, wobei wenigen Nukleinsäure-Sequenzen kein Peptid zugeordnet werden kann und einigen Nukleinsäure-Sequenzen mehr als ein Peptid zugeordnet werden kann. Wie bereits oben erwähnt, sind sowohl die ermittelten Nukleinsäure-Sequenzen, als auch die den Nukleinsäure-Sequenzen zugeordneten Peptid-Sequenzen Gegenstand der vorliegenden Erfindung.

TABELLE I

Sequenz ID No.:	Wahrscheinlichkeit für eine spezifische Expression im Tumorgewebe %	Funktion	Länge des Ausgangs-EST in Basen	Länge der angemeldeten Sequenz in Basen	Chromosomale Lokalisation
1	99.54	Identisch zum humanen IgG aus V-D-J6 Region	202	202	unbekannt
2	99.51	unbekannt	287	1926	unbekannt
3	99.2	unbekannt	196	762	unbekannt
4	98.58	unbekannt	295	918	unbekannt
5	97.43	unbekannt	303	1146	unbekannt
6	97.43	Identisch zum Kaposi Sarcoma-assoziierten Herpesvirus Glycoprotein M	173	2407	
7	95.29	Hyaluroneclin (HN) ist ein Glykoprotein, dass Hyaluron bindet und oft in humanen Tumoren gefunden wird	204	1471	unbekannt
8	95.29	unbekannt	186	1732	unbekannt
9	95.29	unbekannt	197	989	unbekannt
10	91.23	unbekannt	150	150	unbekannt
11	91.23	unbekannt	286	1467	unbekannt
12	91.23	unbekannt	221	895	unbekannt
13	81.23	mitochondriales Enzym	223	467	Chromosom 7
14	81.23	Identisch zum humanen Collagen I (alpha-1 Kette)	198	511	
15	100	Identisch zum humanen Keralin K7 (Typ II)	186	1899	
16	100	unbekannt	228	758	unbekannt
17	99.97	Identisch zum humanen Cofilin	123	302	
18	100	das H19 Gen wird nur vom maternalen Chromosom exprimiert und stellt möglicherweise ein Tumor-Suppressorgen dar	230	824	unbekannt
19	100	Identisch zum humanen IGF-2 Wachstumsfaktor	80	2190	
20	99.56	unbekannt	271	2565	unbekannt
21	99.26	Identisch zum humanen IgV-L (Klon VL 29-1)	227	461	
22	100	Identisch zum humanen H19	213	2086	
23	98.77	mitochondriales Enzym	203	1348	unbekannt
24	100	Identisch zum humanen Anti-Hepatitis A IgG Variable Region	209	358	
25	96.49	Identisch zum humanen Saposin	89	89	
26	94.05	Gen, dass durch IL6 induziert wird	229	1632	unbekannt
27	94.05	unbekannt	195	2872	unbekannt
28	100	Identisch zu Immunoglobulin M schwere Kette V Region	207	496	
29	100	Identisch zur humanen Ig schweren Kette (variable Region)	212	397	
30	100	Identisch zur cDNA, die die leichte Kette eines monoklonalen Antikörpers kodiert, der gegen das humane Cytomegalovirus 65 kD Protein gerichtet ist	243	772	

Sequenz ID No.:	Wahrscheinlichkeit für eine spezifische Expression im Tumorgewebe %	Funktion	Länge des Ausgangs-EST in Basen	Länge der angemeldeten Sequenz in Basen	Chromosomale Lokalisation
31	90.26	unbekannt	210	1031	unbekannt
32	90.26	mitochondriales Enzym	184	739	unbekannt
33	100	Identisch zur Ig kappa leichten Kette variable Region D11.	89	651	
34	99.76	Identisch zum humanen hsp27	197	823	
35	100	Identisch zur Ig schweren Kettevariablen Region V(2-1) (v(h)-iv Familie).	231	457	
36	93.95	unbekannt	203	1203	unbekannt
37	93.95	Identisch zur Immunglobulin leichten Kette variable Region (lambda-IIb Untergruppe) von IgM Rheuma-bezogener Faktoren.	207	207	
38	100	Identisch zu US-Patent	135	346	
39	100	Identisch zu einer Immunglobulin lambda leichten Kette (X57812)	237	926	
40	99.42	Identisch zur humanen hPGI mRNA, die das "bone small proteoglycan I" kodiert	73	2384	
41	99.99	Identisch zu patentierter Sequenz	194	334	
42	99.77	Identisch zu patentierter Sequenz	98	845	
43	99.11	Identisch zur CD24 "signal transducer mRNA"	178	2233	
44	97.85	Identisch zur humanen Ig schweren Kette variablen Region (Klon M49)	242	243	
45	94.61	unbekannt	220	817	unbekannt
46	100	Identisch zur Ig Alpha 1- Alpha 2m=Immunglobulin A1-A2 lambda hybrid GAU schwere Kette (secreted alpha chain)	200	1644	
47	93.7	Identisch zu Pro-alpha 2(I) collagen (COL1A2) Gen	262	1133	
48	99.54	Identisch zu humanen NIC (Natural Inhibitor of Collagenase)	180	969	
49	99.28	Identisch zur humanen Ferritin L Kette	104	617	
50	97.9	Identisch zum humanen Calcyclin Gen (auch Prolactin-Rezeptor assoziiertes Protein)	210	704	

TABELLE II

5

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DNA-Sequenzen Seq. ID. No.	Peptid-Sequenzen (ORF's) Seq. ID. No.
2	128
	129
	130
3	131
	132
	133
4	134
	135
	136
5	137
	138
	139
7	140
	141
	142
8	143
9	144
	145
	146
10	147
	148
	149
11	150
	151
	152
12	153
	154
	155
13	156
	157
	158
16	159
	160
	161

DNA-Sequenzen Seq. ID. No.	Peptid-Sequenzen (ORF's) Seq. ID. No.
18	162
	163
	164
	165
20	166
	167
	168
23	169
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26	171
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27	173
31	174
32	175
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36	178
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45	181
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Die erfinderischen Nukleinsäure-Sequenzen Seq. ID No. 1 bis Seq. ID No. 50 der ermittelten Kandidatengene und die ermittelten Aminosäure-Sequenzen Seq. ID No. 51–106 werden in dem nachfolgenden Sequenzprotokoll beschrieben.

Sequenzprotokoll

(1) ALLGEMEINE INFORMATION:

(i) ANMELDER:

(A) NAME: metaGen - Gesellschaft für Genomforschung mbH

(B) STRASSE: Ihnestrasse 63

(C) STADT: Berlin

(E) LAND: Deutschland

(F) POST CODE (ZIP): D-14195

(G) TELEFON: (030)-8413 1673

(H) TELEFAX: (030)-8413 1674

(ii) TITEL DER ERFINDUNG: Menschliche Nukleinsäure-Sequenzen aus
Blasentumorgewebe

(iii) Anzahl der Sequenzen: 106

(iv) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy disk

(B) COMPUTER: IBM PC compatible

(C) OPERATING SYSTEM: PC-DOS/MS-DOS

(D) SOFTWARE: Patentln Release #1.0, Version #1.25 (EPO)

(2) INFORMATION ÜBER SEQ ID NO: 1:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 202 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 1

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tgagagtcacat ggacctcctg cacaagaaca tgaacacacct gtgggtttcttc ctccctcctgg 60
tggcagctcc _cagatgggtc ctgtcccagg tgcagctaca gcagtggggc gcaggactgt120
tgaagccttc ggagaccctg tccctaacat gcgctgtctc cggtagactct tccagtactt180
actactggga ttggatccgc ca 202
    
```

(2) INFORMATION ÜBER SEQ ID NO: 2:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1926 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 2

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ttgcgatggc tgaatggactg ttgctctcta accaaaggac cctagcgggc tcaacaattg 60
tcaagagcag ttggtggttc tgaatacaat cctcagccaa ggatccctcc tgtgttacag 120
atggatcagc taaaacaagc caaactgaa gacacaaaga atgaggtag gttcattgaa 180
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gcccagacat aggatgggtc agtgggcatg gctctattcc tattctcaa ccatgccagt 600
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acaactgacc tgtctccttc acatagtcca tatcaccaca aatcacacaa caaaaaggag1320
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attattatat tcatatctct acgtgcgaaa ctttctgcct caatgtttac tgtgcctttg1860
tttttgctag tgtgtgttgt tgaaaaaaa aacattccct gcctaagtta gttttggcaa1920
agtatt

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(2) INFORMATION ÜBER SEQ ID NO: 3:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 762 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 3

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ctccactgca accacccaga gccatggctc cccgaggctg catcgtagct gtctttgcca 60
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acctgatgct gtgccagcca cacaagagat gtggggacaa gttctacgac cccctgcagc180
actgttgcta tgatgatgcc gtcgtgccct tggccaggac ccagacgtgt ggaaactgca240
ccttcagagt ctgctttgag cagtgtgcc cctggacctt catggtgaag ctgataaacc300
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ggtgttacct gagatctggg atgctgagtg gctgtttggg ggccagagaa acacacactc480
aactgccac ttcattctgt gacctgtctg aggccacccc tgcgctgcc ctgaggaggc540
ccacaggtcc ccttctagaa ttctggacag catgagatgc gtgtgctgat gggggcccag600
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caggagcaaa gcacaggatc ataataaatt tatgtacttt aa 762

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(2) INFORMATION ÜBER SEQ ID NO: 4:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 918 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 4

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ctcgagccgc tcgagccgat tcggctcgag ccttcccgtc ccctgcttgc aaagtgggtg 60
tgccccaagg tccgctcca gccacgtgg gtgctgctgg ccaagcttc ccttcctttg120
agagaggttt ccgctgtagg agcagagctt ccgggctgcg ctcttcgttg cccagtttcc180

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getcagtggt cgcgtctccg cccccaccc accagtcacg ctgcattctc ggcggggctc240
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ccgagaagcg agagaagctg gcgaacggcg gggagggaat ggaggaggcg accgttgta360
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gaagggggccc ccacgcaaac tcaaattccc tgagcctcaa gaggtggtgg aagagttgaa600
gaagtacctg tcgtaggag atttgggtag aagccctcat gctgagcttt gtgtccctgg660
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gtttcttccc tgccagaaat gaaggttcag ttatgaggca accctctagt aaggcattgt780
aaaagtact ggatttggtt taataaaagt tgaaataaag taaaagaaaa aaaaaaaga840
gaaaaaagaa aaagaaaaaa agaaagaaga aaaaagaaag gagaagcgag agaaaggag900
gccgcggggc gcggcggc 918

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(2) INFORMATION ÜBER SEQ ID NO: 5:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1146 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 5

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tcagtttagt ggcagggtgg ttttttaatt ttcttctgtg gctggatttt tggtgtttgt 60
ttaaataact cttctgggaa gttggtttat aagcctttgc cagggtgaac tggtgtgaaa 120
taccaccac taaagttttt taagtccat attttctcca ttttgccttc ttatgtattt 180
tcaagattat tctgtgcact ttaaatttac ttaacttacc ataatgcag tgtgactttt 240
cccacacact ggattgtgag gctcttaact tcttaaaagt ataatggcat cttgtgaatc 300
ctataagcag tctttatgtc tcttaacatt cacacctact ttttaaaaac aaatattatt 360
actattttta ttattgtttg tcctttataa attttcttaa agattaagaa aatttaagac 420
cccattgagt tactgtaatg caattcaact ttgagttatc ttttaaatat gtcttgata 480
gttcataatc atggctgaaa cttgaccaca ctattgctga ttgtatggtt ttcacctgga 540
caccgtgtag aatgcttgat tacttgtagt cttcttatgc taatatgtc tgggctggag 600
aaatgaaatc ctcaagccat caggatttgc tatttaagt gcttgacaac tgggccacca 660
aagaacttga acttcacctt ttaggatttg agctgttctg gaacacattg ctgcactttg 720
gaaagtcaaa atcaagtgcc agtggcgccc tttccataga gaatttgccc agctttgctt 780
taaaagatgt cttgtttttt atatacacat aatcaatagg tccaatctgc tctcaaggcc 840
ttggtccttg tgggattcct tcaccaatta cttaatttaa aaatggctgc aactgtaaga 900
acccttgctc gatataattg caactatgct cccatttaca aatgtacett ctaatgtca 960
gttgccagggt tccaatgcaa aggtggcggt gactcccttt gtgtgggtgg gggttgtggg1020
tagtggtgaa ggaccgatat cagaaaaatg ccttcaagtg tactaattta ttaataaaca1080

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ttaggtgttt gttaaaaaaa aaaaaaaaaa aaaaaaaazag, ggtgggtsgt gttggtgtgtall140
ttggtt 1146

5 (2) INFORMATION ÜBER SEQ ID NO: 6:

(i) SEQUENZ CHARAKTERISTIK:

- 10 (A) LÄNGE: 2407 Basenpaare
(B) TYP: Nukleinsäure
(C) STRANG: einzel
(D) TOPOLOGIE: linear

15 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

20 (iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

25 (vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
(C) ORGAN:

30 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 6

35
gagtgaagtga gtgtgttgca tcgaattaag gactcttgaa gagaagagag gtccattcag 60
ggttggtccag attgaagtga ggtctcacgg tgaaaagaaa aggaaaatat tcagactctc 120
ttgaaatcca aagagcaaga agtaaatgaa cttctgcaaa aattccagca agctcaggaa 180
40 gaacttgagcag aaatgaaaag atactctgag agctcttcaa aactggagga agataaagat 240
aaaaagataa atgagatgtc gaaggaagtc accaaattga aggaggcctt gaacagcctc 300
tcccagctct cctactcaac aaagctcatcc aaaaggcaga gtcagcagct ggaggcgctg 360
cagcagcaag tcaaacagct ccagaaccag ctggcggaat gcaagaaaca acaccaggag 420
gtcatatcag tttaacagaat gcatactctg tatgctgtgc agggccagat ggatgaagat 480
45 gtccagaaaag tactgaagca aatccttacc atgtgtaaaa accagtctca aaagaagtaa 540
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60 tacagccaaa agaatgcct catagtctctt aacctcaact tttgtagaag tatttttttc1380
tctgtaatat ttttattggc tcataaagat gttttcatat ctgaactcct aaataagtgal1440
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 cgagtactct ggttaaatat tgaaaagtta tatgctgtag tttttagtag tttgtctttg 2340
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 atgaaaa 2407

(2) INFORMATION ÜBER SEQ ID NO: 7:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 1471 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 7

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 gccctcggg tccgacctt tgagcgttct gtcctggcgc cagctacctc gctcctcggc 120
 gccatgacca caaccaccac cttcaaggga gtgcaccca acagcaggaa tagctcccga 180
 gttttgcggc ctccaggttg tggatccaat ttttcattag gttttgatga accaacagaa 240
 caacctgtga ggaagaacaa aatggcctct aatatctttg ggacacctga agaaaaataa 300
 gcttcttggt ccaagtcagc aggtgccaag tctagtggg gcagggaga cttggagtca 360
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 aacctgatgc tgtgtgcatg atgctgggtg ttgaccatga aaggaaagtc tcattccttaa 1380

aatgtgttgt acttcacaat cctggactgt tgcttcaagt aaaccaatc cccattttgal1440
 aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa a 1471

5 (2) INFORMATION ÜBER SEQ ID NO: 8:

(i) SEQUENZ CHARAKTERISTIK:

- 10 (A) LÄNGE: 1732 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

- 15 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

- 20 (iii) HYPOTHETISCH: NEIN

- (iii) ANTI-SENSE: NEIN

25 (vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

30 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

35 (xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 8

gcagaaccta cgctgacgg gccggcgcc ggctgagccg cgctgocgag cgacgcggga 60
 atgaagcggg cgctgggag gcgaaagggc gtgtggttg gcctgaggaa gatacttttc 120
 40 tgtgttttgg ggtgtgacat tgccattcca tttctcatca aactatgtcc tggaaacag 180
 gccaaactga ttttcttgaa tttcgtgaaga gttccctatt tcattgattt gaaaaacca 240
 caggatcaag gtttgaatca cacgtgtaac tactacctgc agccagagga agacgtgacc 300
 attggagtct ggcacaccgt cctgcagtc tggtggaaga acgccaagg caaagaccag 360
 atgtggtatg aggatgcctt ggcttcagc caccctatca ttctgtacct gcatgggaac 420
 45 gcaggtacca gaggaggcga ccaccgcgtg gagctttaca aggtgctgag ttcccttggg 480
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 cggggcatga cctatgacgc actccacgtt tttgactgga tcaaagcaag aagtggtgac 600
 aaccccggtg acatctgggg ccactctctg ggcactggcg tggcgacaaa tctggtgagg 660
 cgctctgtg agcgagagac gccctccagat gcccttatat tggaaatctcc attcactaat 720
 50 atccgtgaag aagctaagag ccatccattt tcagtatat atcgatactt ccctgggttt 780
 gactggttct tccttgatcc tattacaagt agtggaatta aatttgcaaa tgatgaaac 840
 gtgaagcaca tctcctgtcc cctgtcatc ctgcacgtg aggacgacc ggtggtgccc 900
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 55 aaagttcagt ttgtgccctt tcattcagac cttggctaca ggcacaaata catttacaag1020
 agccctgagc tgccacggat actgagggaa ttcctgggga agtcggagcc tgagcaccag1080
 cactgagcct ggccgtggga aggaagcatg aagacctctg ccctcctccc gtttccctcc1140
 agtcagcagc ccggtatcct gaagccccgg ggggcccggc cctgcaatgc tcaggagccc1200
 agctcgcacc tggagagcac ctcatatccc aggcggggag gccctgagc gcctgagtg1260
 60 cccggaggcc tgagcatggc tgtgtggaag cgtgggtgg caggcatgtg gctctccttg1320
 ccgcccctoa acctgagatc ttgttgggag acttaatggc agcaggcagc catcactgcc1380
 tgggtgatgc tgactgagc tggacagggg gaggccgggc aggggactct tggggctcgg1440
 gaccatgctg agcttttttg caccacccac agagaacgtg ggggccaggt tctttctgca1500
 ccttcccagc acatgcagaa tgactccagt ggttccatcg tcccctcctg ccctgtgtac1560
 65 ctgcttgcc tttctagctg cccacacctc cctgggctgg cccactcacc cacagtggaa1620
 gtgcccggga tctgcacttc ctcccctttc acctacctgt acacctaacc tggccttaga1680
 ctgagcttta tttaagaata aaatcgtggt ggtggtcaaa aagacactct gc 1732

(2) INFORMATION ÜBER SEQ ID NO: 9:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 989 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 9

```

cggtctgagc gtgatcgctg actcagctga ccctgcggga ccggaaaag aaattcccgg 60
gccctggctt cttggcgcca tgaggttccg gttctgtggt gatctggact gtcccactg120
ggtcctggca gaaatcagca cgctggccaa gatgtcctct gtgaagttgc ggctgctctg180
caccaggtac taaaggagct gctgggacag gggattgatt atgagaagat cctgaagctc240
acggctgacg ccaagtttga gtcaggcgat gtgaaggcca cagtggcagt gctgagtttc300
atcctctcca gtgcggccaa gcacagtgtc gatggcgaat ccttgtccag tgaactgcag360
cagctggggc tgcccaaaga gcacgcggcc agcctgtgcc gctgttatga ggagaagcaa420
agcccccttc agaagcactt gcgggtctgc agcctacgca tgaatagggt ggacaggtgtg480
ggctggcggg tggactacac cctgagctcc agcctgctgc aatccgtgga agagcccatg540
gtgcacctgc ggctggaggt ggcagctgcc ccagggaccc cagcccagcc tgttgccatg600
tccctctcag cagacaagtt ccaggtcctc ctggcagaac tgaagcaggc ccagaccctg660
atgagctccc tgggctgagg agaaggggtg tccaggcctg tgtggagccg ccctgcccgt720
atggagtca c gccctctgaa ctgctcttcg ggaggcagcc ctggttctag gatgctgagg780
ccctggccc gactctggcc tcccagatcc ccagctgcct cacttctctc ttgagaactt840
ggctcagggc tcttgaggac ctttcccagc attaccttcc cttcccttga aaggcaattg900
ttggctgttt tcataagcag gaaaaataaa cagaagtata aaagagaaaa aaaaaaaaaa960
aaaaaaaaa aaaaaaaaaa aaaaaaaaaa
989

```

(2) INFORMATION ÜBER SEQ ID NO: 10:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 150 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 10

```

attttatgaa gttgaaaaat agctcacttt aaagctagtt ttgaagacgt gcagctgtga 60
cttgggtctg gttgggggtg ttgtgttttg agtcagccgt tttcactccc actgaggttg120
tcagaacatg cagattgctt cgattttctc
150

```

(2) INFORMATION ÜBER SEQ ID NO: 11:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1467 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 11

```

cgaggaccgg ccttgcgagc ggcgacgact ataaaatggc gcgtgctgca acccgcgccc 60
gcttcggaga gagaaatgct ggggtgcagc ttcaagctta ggaccaccca ccatgcctat 120
ccaggtgctg aagggcctga ccatcactca ttaagaacag aggaggetgc ctgttactcc 180
tggtgttgca tccctccaga cactctgctg tttcctgcct aggcgtggct gcagccatgg 240
ctaggaaagc gctgccaccc acccaacctg gccagagctg gttctgctcc tgetgcaggg 300
60 acaactgagot ggctatctcg gcgcttcggg caagaactgc aacaggctct cctgggtcct 360
gcaggtgtac agccggggccc ctgccttgct cctcagctct cgagagctgc tgetgccggg 420
tgacctgac caacctgata agtgccatc ttcagctacc actgcaaggc cctgagggca 480
acagcagcac ggcactgccc acccggtctg tgatggcctg gtgccagctg ggagtccctc 540
cggcacttcg aggccactga gccacccttc cagccccagc ccaccatgga caggggtatc 600
65 cagcttcctc ctcaacctcg tcctctgccc ctgagccagt gacgccaag gacatgcctg 660
ttaccaggt cctgtaccag cactagctgg tcaagggcat gacagtgtg gaggccgtct 720
tgagatcca ggccatcact ggcagcaggc tgctctccat ggtgccaggg cccgccaggc 780
caccaggctc atgctgggac ccaaccctg gcacaaggac ttggctgctg agccacacac 840

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ccaggagaag gtggataagt gggctaccaa gggcttcttg caggctaggg gaggaaccac 900
 ccccgcttcc ctattgtgac caggcctatg gggagagctt gtccatagc caccgtgaga 960
 cctgggcttg gctctcaagg acagacaccg cctggcctgg tgctccaggg gtgaagcagg1020
 ccagaatcct gggggagctg ctcttggttt gagctgcatt caggaaagtgc gggacatggt1080 5
 aggggaggca aaaagccttg ggcactaccc tccctgtgga gctgttcggg gtccgtcgag1140
 ctayccacac cctgacacca tgttcaaggg taccggaaga gaagggtgtc tgccccaac1200
 ctcccctgtg ggtgtcactg gccagatgtc atgagggaag caggccttgt gagtggacac1260
 tgaccatgag tccctggggg gagtgatccc ccaggcatcg tgtgccatgt tgcacttctg1320
 cccaggcagc aggggtgggtg ggtaccatgg gtgcccaccc ctccaccaca tggggcccca1380 10
 aagcactgca ggccaagcag ggcaacccca cacccttgac ataaaagcat cttgaagctt1440
 ttaaaaaaaa aaaaaaaaaa aaaataa 1467

(2) INFORMATION ÜBER SEQ ID NO: 12:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 895 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 12

ctcgagccgc tcgagaacct cacttcctta ctctccaaa aagaagtggg gaaagaacca 60
 tcaaaccttt cctcctgact taccaaacca ggaaacagc aggagagggt ggctcaggac120
 ttagggacag ggtatagctt agatgggtga aagcaaaggg gagcaggaag ttgtaaatca180
 ctggctaagt gaaaaaggag acagctaact ctaggatgaa gctgtgacta ggctggagtt240
 gcttccttga agatgggact ccttgggtat caagacctat gccacatcac actggggcta300 50
 gggaagtagg tgatgccagc cctcaagtct gtcttcagcc agggacttga gaagtatat360
 tgggcagtgg ctccaatctg tggaccagta tttcagcttt ccctgaagat caggcagggt420
 gccattcatt gtctttctct cctagccccc tcaggaaaga aggactatat ttgtactgta480
 ccctaggggt tctggaaggg aaaacatgga atcaggattc tatagactga taggccctat540
 ccacaagggc catgactggg aaaaggtagg ggagcagaag gagaattggg attttaggg600 55
 gcagtacgct caccctaao ttttgggtgg ctggggcatg tcttgaggcc cagactgtta660
 agcaggctct gctggcctgt ttactcgtca ccacctctgc acctgctgtc ttgagactcc720
 atccagcccc aggcacgcca cctgtcctg agcctccact atctccctgt gacgggtgaa780
 ctctgtgtac tgtgtctcgg gtccatatat gaattgtgag cagggttcat ctattttaa840 60
 cacagatgtt taaaaataa agattatttc aaacaaaaa gaaaaaaa aaaaa 895

(2) INFORMATION ÜBER SEQ ID NO: 13:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 467 Basenpaare

- (B) TYP: Nukleinsäure
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

- (vi) HERKUNFT:
(A) ORGANISMUS: MENSCH
(C) ORGAN:

(vii) SONSTIGE HERKUNFT:
(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 13

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accagcagc cctcgcgcgg tccggcacag cggacaccag gactccaaaa tggcgctcagt 60
tgtaccagtg aaggacaaga aacttctgga ggtcaaactg ggggagctgc caagctggat120
cttgatgcgg gacttcagtc ctagtggcat ttctggagcg ttcaaagag gttactaccg180
gtactacaac aagtacatca atgtgaagaa ggggagcatc tcggggatta ccatgggtgct240
ggcatgctac gtgctcttta gctactcctt ttctacaag catctcaagc acgagcggct300
ccgcaaatac cactgaagag gacacactct gcaccccccc accccacgac cttggcccga360
gcccctccgt gaggaacaca atctcaatcg ttgctgaatc ctttcatatc ctaataggaa420
ttaacctcca aataaaacat gactggtacg tgtaaaaaaa aaaaaaa 467
    
```

(2) INFORMATION ÜBER SEQ ID NO: 14:

- (i) SEQUENZ CHARAKTERISTIK:
(A) LÄNGE: 511 Basenpaare
(B) TYP: Nukleinsäure
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

- (vi) HERKUNFT:
(A) ORGANISMUS: MENSCH
(C) ORGAN:

(vii) SONSTIGE HERKUNFT:
(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 14

actagttcta gatcgcgagc ggccgcctt tttgttcaaa gtctatTTTT attctctttt 60
 tctttttttt ttttttgtgg atggggactt gtgaattttt ctaaagggtgc tatttaacat120
 gggaggagag cgtgtgcggc tccagccag cccgtgctc actttccacc ctctctccac180
 ctgctcttgg cttctcaggc ctctgctctc cgacctctct cctctgaaac cctcctccac240
 agctgcagcc catcctcccg ggcctctcct agtctgtcct gcgctcctctg tccccgggtt300
 tcagagacaa cttcccaaag cacaagcag tttttccccc taggggtggg aggaagcaaa360
 agactctgta cctattttgt atgtgtataa taatttgaga tgtttttaat tattttgatt420
 gctggaataa agcatgtgga aatgaccca aacataaaaa aaaaaaaaaa aaaaaaaaaa480
 aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa a 511

(2) INFORMATION ÜBER SEQ ID NO: 15:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1899 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 15

tctccaccct ccccttcctt ctttcttttt ccccttcttg cactggatc actcaggcct 60
 cagaatgagg ctgctttatt ggaagctatt ctgacatcac ttccagact gtctcactgt 120
 cttgggacca ggcattggag gcagggttgg gaatcttctt gtgattgttg gtggtggctg 180
 gaggagtga gtggtggag gcggctcagt cgcgggcaact cctgcgactg gcggatgcgg 240
 tccggatgga ataagccttc aggaagccag gacccgcaact gctggagaag ctccaggcat 300
 tgctgcccat ggttcccccg agggtcagcc caatgccacc gccactgcta ctgccaccag 360
 tggaattcat cacagagata ttacaggctc ccaactccatc tccagccaac cggctctcct 420
 cgccctccag cagcttgagg taggtggcga tctcgatgtc cagggccagc ttcacgctca 480
 tgagttcctg gtactcacgc agctgccgtg cdatatcctg cttggcccgc tgcagggcgg 540
 cttccagctc ctctgtcttg gcacgagcat ccttgagcgc cagctcccca cgctcctcag 600
 cctcggaat ggcggcctcc aacttggcac gctggttctt gatgttgctg atctcagcct 660
 gcagcctctg gatggcccgg ttcatctctg aaatctcatt ccgggtattc cggagggtcgt 720
 cccatgctt cccagccttg gcttgagggt tctcaaaact ggtctggtac caggcttcag 780
 cctcagcccg gctgcatctg gccatctcct catactgcgc cttgacctca gcgatgatgc 840
 cgtccaggctc caggagcgga ctgttgctca tggacagcac cacagatgtg tcggagatct 900
 gggactgcag ctctgtcaac tccgtctcat tgagggtcct gaggaagttg atctcatcat 960
 tcagggcac cacttgccc tccagctcca ccttgctcat gtaggcagca tccacatcct1020
 tcttcagcac cacaactca ttctcagcag ctgtgcggcg gtttaatttca tcttcgtact1080
 tattcttgaa gtccctccacc acatcctgcc atctccgcag ctccgcctcc aggcggcccc1140
 catccacctg cagtgcctca agctgacccc gaaggccagc aatctgggcc tcaaagatgt1200
 ctgggaggcg gctgctcttg gccgacttct gctcctgcag cagcgtccac ttggtctcca1260

gcagcttggt ctgctgctcc agaaaccgca ccttytcgat gaagvaggaa aacttytgyt 1320
 tgagggtctt gatctgctcg ctctctcct ggccgaccgg cgggagggag gggtcggcgt 1380
 ccagccgcag cggggccagc aggtcttggt taatggtgac ctgcggtatg ccggcgccca 1440
 5 ccggggcccc ataggcagag cgcacggcca cgcgcgccg tgaggcgccg aggccgtaga 1500
 ggctgctgct gccaaggccg cggggcgag cggagctcag gcgcacctg gcccgcgcc 1560
 ccgagaaggc ggtgagcgc gaggtgaata cggggagct gaagtggatg gacatggtgg 1620
 ctggggccgg atggacctag cggcgggcga ggaggagcgc gactcgtg acctcgggga 1680
 10 cactccgcac ctttatccg cgggagccgg tgctgggctt ccacaggtag ggccggggct 1740
 ggcccggggc accgtttctc tgctgccagg cccctcctgc gcgtccgtcc gccctctgcc 1800
 cgcccgcccc gccgaagccc aggttttcag tccaagcagg gatggtccgg agtaggcagg 1860
 agcgccatcc ctagacggcc gcagagaaca gcgggggac 1899

15 (2) INFORMATION ÜBER SEQ ID NO: 16:

(i) SEQUENZ CHARAKTERISTIK:

- 20 (A) LÄNGE: 758 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

- 25 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

- 30 (iii) HYPOTHETISCH: NEIN

- (iii) ANTI-SENSE: NEIN

35 (vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

40 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 16

45 cggtcgcagc ggctcgagg gcaagaagaa catcacgtgc tgtgacaccg acttgtgcaa 60
 cgccagcggg gcccatgccc tgcagccggc tgcggccatc cttgcgctgc tccctgcact 120
 cggcctgctg ctctggggac cggccagct ataggctctg gggggcccg ctgcagccca 180
 50 cactgggtgt ggtgccccag gctctgtgc cactcctcac agacctggcc cagtgggagc 240
 ctgtcctggt tcctgaggca catcctaacg caagtctgac catgtatgtc tgcaccctg 300
 tgccccaccc tgacctccc atggccctct ctaggactcc cacccgccag atcagctcta 360
 gtgacacaga tccgcctgca gatggccct ccaacctct ctgctgctgt ttccatggcc 420
 cagcattctc cacccttaac cctgtgtca ggcacctct cccccaggaa gccttcctg 480
 55 cccaccccat ctatgacttg agccaggtct ggtccgtggt gtccccgca cccagcagg 540
 gacaggcact caggaggcc cagtaaaggc tgagatgaag tggactgagt agaactggag 600
 gacaagagtc gacgtgagtb cctgggagtc tccagagatg gggcctggag gcctggagga 660
 aggggcccag cctcacattc gtgggctcc ctgaatggca gcctgagcac agcgtaggcc 720
 60 cttataaac acctgttgga taagccaaaa aaaaaaaa 758

(2) INFORMATION ÜBER SEQ ID NO: 17:

65 (i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 302 Basenpaare
 (B) TYP: Nukleinsäure

(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH
(C) ORGAN:

(vii) SONSTIGE HERKUNFT:
(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 17

```
cggtctgacg gtctcgagat caagcatgaa ttgcaagcaa actgctacga ggaggtcaag 60
gaccgctgca ccctggcaga gaagctgggg ggcagtgccg tcctctccct ggagggcaag120
cctttgtgag cccctttctg cgecccttg cctgggagca tctgggcagg cccaacacc180
ttgccctttg ggggtttgca gggctcgccc cctttcctgg ccagaaccgg gagggggctg240
gggggggatt cccaggcagg gggggagggg ccaattccct ttccaacccc caggttgggc300
ca                                     302
```

(2) INFORMATION ÜBER SEQ ID NO: 18:

(i) SEQUENZ CHARAKTERISTIK:
(A) LÄNGE: 824 Basenpaare
(B) TYP: Nukleinsäure
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH
(C) ORGAN:

(vii) SONSTIGE HERKUNFT:
(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 18

```
ggcgaggaca gaggaggcgc gtccggcctt cctgaacacc ttaggctggt ggggctgcgg 60
caagaagcgg gtctgtttct ttacttcttc cactggagtcg gcacactatg gctgccctg120
```

ggctcccaga acccacaaca tgaagaaat ggtgctaccc agctcaagnc tgggccccttg 180
 aatccggaca caaaaccctc tagcttgga atgaatagc tgcacctta aaccactgca 240
 ctacctgact caggaatcgg ctctggaagg tgaagctaga ggaaccagac ctcatcagcc 300
 caacatcaaa gacaccatcg gaacagcagc gcccgagca cccaccccg accggcgact 360
 ccatcttcat ggccaccccc tgcggcgagc ggttgaccac cagccaccac atcatcccag 420
 agctgagctc ctccagcggg atgacgcgt cccaccacc tccctcttct tctttttcat 480
 ccttctgtct ctttgtttct gagctttcct gtctttcctt ttttctgaga gattcaaagc 540
 ctccacgact ctgtttcccc cgtcccttct gaatttaatt tgcactaagt catttgact 600
 ggttgaggtt gtggagacgg ccttgagctc cagtacgagt gtgcgtgagt gtgagccacc 660
 ttggcaagtg cctgtgcagg gcccgccgc cctccatctg ggccgggtga ctggggcgcc 720
 ggctgtgtgc ccgaaggcct caccctggcc ctccggcctt agtctggga gggtccgaac 780
 cgaacatcaa gggaggcaag cctttcaagg catttcatt aatt 824

(2) INFORMATION ÜBER SEQ ID NO: 19:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 2190 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 19

tttttttttt tttttttttt tttttttttt tttttttttt tttcatgact tttaatgctt 60
 tattgggatt gcaagcgtta caaggttaaa gacaaaaccc aagcatggga ttttgccgga 120
 aatatttagc ttaaaggagc tgagttgagt caaacacggg ccgcaagggt gaccgaggcg 180
 gcaggcacag gtgacattca gtgtttggcg tgggggtcct caggtgatgg cagaggaggg 240
 gacccaagag ggggcccccc actgaagaca ttggggacac ggggaggaga caagatggag 300
 agccacgact aggcacggag gtcagacagg cagcccgggc caggatgggt agtgcccag 360
 gggagagctg caaacctggg gacgcaaggg gctggtcggc aagtgcctcc gggaacaccc 420
 actccggcga ggcagaatat aacactgggt ggggtgggtg cctgacgaat gggcaggtaa 480
 tttggggtgc ctgcaagcgt tttggatctc aggccaatgt gggttccaca attgtgacaa 540
 tttgctctt tgggctctg tccaatgttc cgaatggccc actcacaggg cgcttgccga 600
 gggaccctct gcgacgctc agctcgagcc gaaatgaggg aacccccaaa tttcatgtca 660
 attgatctat tccccctt tgtttcttgg ggcagttttt tttttacccc tccttagctt 720
 tatgcgctca gaaaccaaata taaaccccc ccccatgtaa caggggggca gtgacaaaag 780
 caagaacgca cgaagccagc ctggagacca ccacgtcctg ccccccgcca tttatcgccc 840
 tgattggatt ttgtttttca tctgtccctg ttgcttgggt tgagttgagg gtggagcctc 900
 ctggggggca ctggccactg agccccctt gagaagttag aggggagtgg agaaggccac 960
 tgtccggcct ggcttctggg gacagtggct ggtccccaga agtcctgagg gcggaggggg 1020
 ggggtgggca ggggtctcct aggtgtcagg aggggtgctg gaggccacag gagggggctc 1080
 ctggctggcc tgaggctggc cggaggggaa ggggctagca ggtgtgtaaa cagagggttc 1140

catcaggctg gggcaggggt gccgccttcc gcacacttga ggaacccttc cccctccctc 120
 ggtgacatct tgcccccccc tcagcaccct gccttgcttc caggaggtcc gaagctctgt 1260
 gggacctctt gggggcaagg tggggtgagg ccggggagta gggaggtcag gcgggtctga 1320
 gccacagag caggagagct gccaggtctg cccatcgacc aggttgcttg ggccccggag 1380
 cccacgggtc tggtagtgcc atagcagcca ccaccgccc gcctagggct gcggcaggga 1440
 ctcggcctct gggaggttta cctcgcccc acttggtccc ccagctcagc cccctgcac 1500
 gcagcccgac tagcagtcta gaggcctgag gcttctgggt cctggtgacg gggctggcat 1560
 gaccccgggg gtggtccatg ccagtcggcc tcagtcgag agggtccttc ggcaagcgcc 1620
 ctgtgagtg gcatctcgga acattggaca gaagccaaa gagccaaatt gtcacaattg 1680
 tggaaccac attggcctga gatccaaaac gcttcgaggc accccaaatt acctgcccc 1740
 tcgtcaggac acccaccac ccagtggtat attctgcctc gccggagtgg gtgttcccc 1800
 gggcacttgc cgaccagccc cttgcgtccc cagggttgca gctctccctt gggccactaa 1860
 ccatctggc ccgggtgccc tgtctgacct ccgtgcctag tcgtggctct ccatcttgct 1920
 tcctcccggt gtccccaatg tcttcagtg ggggccccct cttgggtccc ctcctctgcc 1980
 atcacctgaa gacccccacg ccaaacactg aatgtcacct gtgctgccg cctcggtcca 2040
 ccttgccggc cgtgtttgac tcaactcagc tcctttaacg ctaatatctt cggcaaaatc 2100
 ccatgcttgg gttttgtctt taaccttgta acgcttgcaa tccaataaa gcattaaaag 2160
 tcctgaaatt caaaaaaaaa aaaaaaaaaa 2190

(2) INFORMATION ÜBER SEQ ID NO: 20:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 2565 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 20

ctccccacc tgtggccgc aagccgtctg tgggagtcgc ggaccccgcc tccccagtt 60
 accctcgagc tgagccctt actgctctc ccaccaatgg gctccctcac acccaggaca 120
 ggactaagag ggagctggcg gagaatggag gtgtcctgca gctggtgggc ccagaggaga 180
 agatgggctt cccgggtca gactcacaga aagagctggc ctgaccacca ggcacctcac 240
 tggcactgct gacccatccc agaaacacaa tctcagggac ccgagcagct ccaaggacga 300
 gaggatacag cagacacaac ctaatagaga gggcgccctg agccttaacc tccacggcct 360
 tcgatactta tgcaagcctg gtgttgctcc tgtcctcaga gtcactctgc gctcatgctt 420
 tttcccgaaat ggggtcacct ctggcagttg ccgcttcagt cttggcctta gcctcatctt 480
 gaagtgggta gctggcggga gaggtggct gcgccccctg ctggccctga ggctgcagag 540
 ttgggagcag gacacctcac ctgagtttca ttttttttca tgtccaaacc atgcacatac 600
 tatagtccag aatcaaagca cttttgaaaa gtggctgcat ggccatcctc caggggccag 660
 gaagtgtcat tccaaggggc tgtttacatg gcagcagaat ccatccccgg cagtcagccc 720
 atagcttggg accagctctgt gccctcctgc ccagtcagct ttactcctct tggttcctga 780
 aggtggccaa gtcattgtgt tcccacaggc ttctctaggc tgggggcagg tgtggggctg 840

5 tgggaattcca aagcacaaaa ggtgcagagg ggattggcct tectgtgcct caactcacc 900
 accaccctcc tgccttccag ttctgccagg tgctccatg tggggacaag taggagactg 960
 ccagggccca aagaaatggg tgagcagtag agtcatctcg gggcacttgg cagtgtcaag 1020
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 aggagccttt gggctgagca agcactgagg aggtggatgg aaggagcat ctggagggg 1140
 ggagcttccct tgagcagtgg gccagggcct ggccctccac acttcattct ctgaccttcc 1200
 tctctectca ttctgggtgca tgtcctttct gcagctgcct ttcagcacag gtggttccac 1260
 tgggggcagc taacgctgag tgacaaggat gggaagccac aggtgcattt tactcaagtc 1320
 10 ttctctagtc aatgaggggc acccagtgtc tctagggcag gctgggtggg ggtcccctag 1380
 gtatcagcct ctcttactgt actctccggg aatgttaacc tttctatttt cagcctgtgc 1440
 cacctgtcta ggcaagctgg cttccccatt ggccccctgt ggtccacagc agcgtggctg 1500
 cccccaggg ccaccgcttc tttcttgatc ctctttcctt aacagtgaact tgggcttgag 1560
 tctggcaagg aaccttgctt ttgacttcac caccaaggag agaggttgac atgacctccc 1620
 15 cgccccctca ccaaggtggg gaacagaggg gatgtgggtg gagccagggt cctctggccc 1680
 tctccagggt gttttccact agtcaactact gtcttctcct tgtagctaata caatcaatat 1740
 tcttcccttg cctgtgggca gtggagagtg ctgctgggtg tacgctgcac ctgcccactg 1800
 agttggggaa agaggataat cagtgagcac tgttctgctc agagctcctg atctacccca 1860
 20 ccccttagga tccaggactg ggtcaaagct gcatgaaacc aggccctggc agcaacctgg 1920
 gaatggctgg aggtgggaga gaacctgact tctctttccc tctccctcct ccaacattac 1980
 tgggaactcta tctgttagg atcttctgag cttgtttccc tgcctgggtg gacagaggac 2040
 aaaggagaag ggaggtgcta gaagaggcag cccttctttg tctctgggg taaatgagct 2100
 tgacctagag taaatggaga gacaaaagc ctctgatttt taatttccat aaaatgttag 2160
 25 aagtatatat atacatatat atatttcttt aaatttttga gtctttgata tgtctaaaaa 2220
 tccattccct ctgccctgaa gctgagtga gacacatgaa gaaaactgtg tttcatttaa 2280
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 agggcagtggt tctgggcccc catttagaag ggaaaatgtt ttgctgctg cacacattgg 2400
 30 acccaagtat gggcctcttc tgccctagtag tgccaaaggg actgttaagg tgtcttgtcc 2460
 atcttctacc cccaccccc cattacaggg taaagggaac cccagactag gtgagggggc 2520
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35 (2) INFORMATION ÜBER SEQ ID NO: 21:

(i) SEQUENZ CHARAKTERISTIK:

- 40 (A) LÄNGE: 461 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

- 45 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

- 50 (iii) HYPOTHETISCH: NEIN

- (iii) ANTI-SENSE: NEIN

55 (vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

60 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 21

65 gttctctcct agcggctcgc tcagaagcag agttctgggg tgtctccacc atggcctgga 60
 cccctctctg gctcactctc ctcactcttt gcatagggtc tgtggtttct tctgagctga 120
 ctcaggaccc tgctgtgtct gtggccttgg gacagacagt caggatcaca tgccaaggag 180

acagcctcag aagctattat gcaagctggt accagcagaa gccaggagag gccctgtac240
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 gctcaggaaa cacagcttcc ttgaccatca ctggggctca ggcggaagat gaggctgact360
 attacttgta aactccccgg gacagcaagt gggtaaccaa tgtgggtatt ccggcgggag420
 ggacccaagc ttgaccggt ctttaaggtca gcccaaagg c 461

(2) INFORMATION ÜBER SEQ ID NO: 22:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 2096 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(II) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 22

atccccgtca cttttggtta caggacgtgg cagctggttg gacgagggga gctggtgggc 60
 agggtttgat cccagggcct gggcaacgga ggtgtagctg gcagcagcgg gcaggtgagg 120
 accccatctg ccgggcaggt gagtcccttc cctccccagg cctcgtcttc ccagccttct 180
 gaaagaagga ggttttaggg atcgagggct ggcggggaga agcagacacc ctcccagcag 240
 agggcagga tggggcagg agagtttagca aaggtgacat cttctcgggg ggagccgaga 300
 ctgcgcaagg ctggggggtt atgggcccgt tccaggcaga aagagcaaga gggcagggag 360
 ggagcacagg ggtggccagc gtaggggtcca gcacgtgggg tggtagccca ggcctgggtc 420
 agacagggac atggcagggg acacaggaca gaggggtccc cagctgccac ctacccacc 480
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 ggagcctcga gggctggaga gcgggaagac aggcagtgtc cggggagttg cagcaggacg 600
 tcaccaggag ggcgaacggc cacgggaggg gggccccggg acattgcgca caaaggaggc 660
 tgcaggggct cggcctgcgg gcgcgggtcc cagcaggcac tgcggcccag ggtctggtgc 720
 ggagagggcc cacagtggac ttggtgacgc tgtatgccct caccgctcag cccctggggc 780
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 gcgagggcgg cggggcgagg tgaatgagct ctcaggaggg aggatggtgc aggcagggg 900
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 gcctcgcggg cggcgacgga gccgggatcg gtgcctcagc gttcgggctg gagacgaggc1020
 caggtctcca gctggggtgg acgtgcccac cagctgccga aggcgaagac ccaggtccgg1080
 tggagctgac aagcaggaca tgacatggtc cggtgtgacg gcgaggacag aggaggcgcg1140
 tccggccttc ctgaacacct taggctgggt gggctgcggc aagaagcggg tctgtttctt1200
 tacttcctcc acggagtccg cacactatgg ctgccctctg ggctcccaga acccacaaca1260
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 ctctggaagg tgaagctaga ggaaccagac ctcatcagcc caacatcaaa gacaccatcg1440
 gaacagcagc gcccgagca cccaccccg accggcgact ccatcttcat ggccaccccc1500
 tgcggcgagc ggttgaccac cagccaccac atcatcccag agctgagctc ctccagcggg1560
 atgacggcgt cccaccacc tccctcttct tctttttcat ccttctgtct ctttgtttct1620
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cgctcccttct gaatttaatt tgcactaagt catttgcact gggttgagtt gtggagacgg1740
 ccttgagtct cagtacgagt gtgcgtgagt gtgagccacc ttggcaagtg cctgtgcagg1800
 gcccgccgc cctccatctg gcccggtga ctggcgccg gctgtgtgcc cgaggcctca1860
 5 cctgcccctc gcctagtctg gaagctccga ccgacatcac ggagcagcct tcaagcattc1920
 cattacgccc catctcgctc tgtgcccctc cccaccaggg cttcagcagg agccctggac1980
 tcatcatcaa taaacactgt tacagcaaaa aaaaaaaga aaaaaaaag aaaagaaaaa2040
 aaagaaaaag aaaaggaaaa aaagagaggg aggaagaagg agaaaagga gtgtgg 2096

10

(2) INFORMATION ÜBER SEQ ID NO: 23:

15

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1348 Basenpaare

(B) TYP: Nukleinsäure

20

(C) STRANG: einzel

(D) TOPOLOGIE: linear

25

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

30

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

35

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

40

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 23

ctctcgcgag gcccagaga gcaggcgtg ggcagtgtg aggtcgttg agtcacttcc 60
 gcgtcaccag ctcctgtgcc tgccagtcgg tgccctccc gctccagcca tgctctccgc 120
 cctcgcccg cctgccagcg ctgctctccg ccgcagcttc agcacctcgg cccagaacaa 180
 tgctaaagta gctgtgctag gggcctctgg aggcacggg cagccacttt cacttctcct 240
 gaagaacagc ccttgggtga gccgcctgac cctctatgat atcgcgaca caccggagt 300
 50 ggccgcagat ctgagccaca tcgagaccaa agccgctgtg aaaggctacc tcggacctga 360
 acagctgcct gactgcctga aaggttgtga tgtggtagt attccggctg gagtccccag 420
 aaagccaggc atgacccggg acgacctgtt caacaccaat gccacgattg tggccaccct 480
 gaccgctgcc tgtgcccgag actgcccggg agccatgac tcggtcattg ccaatccggt 540
 taattccacc atccccatca cagcagaagt ttccaagaag catggagtgt acaaccccaa 600
 55 caaaatcttc ggcgtgacga ccctggacat cgtcagagcc aacacctttg ttgcagagct 660
 gaagggtttg gatccagctc gactcaacgt ccctgtcatt ggtggccatg ctgggaagac 720
 catcatcccc ctgatctctc agtgaccccc caaggtggac tttcccagg accagctgac 780
 agcactcact gggcgatcc aggaggccgg cacggagggt gtcaaggcta aagccggagc 840
 60 aggtctgcc accctctcca tggcgtatgc cggcgccgcg tttgtcttct cccttgtgga 900
 tgcaatgaat ggaaaggaag gtgttgtgga atgttccttc gtttaagtcac aggaacgga 960
 atgtacctac ttctccacac cgctgctgct tgggaaaaag ggcatcgaga agaacctggg1020
 catcggaaca gtctcctctt ttgaggagaa gatgatctcg gatgccatcc ccgagctgaal080
 ggcctccatc aagaaggggg aagatttcgt gaagaccctg aagtgaaggc ctgtgacggg1140
 65 tggccagttt ccttaattta tgaaggcatc atgtcactgc aaagccgttg cagataaact1200
 ttgtatttta atttgctttg gtgatgatta ctgtattgac atcatcatgc cttccaaatt1260
 gtgggtggct ctgtgggcgc atcaataaaa gccgtccttg attttaaaaa aaaaaaaaaa1320
 aaaaaaaaaa aaaaaaaaaa aaaaaaaa 1348

(2) INFORMATION ÜBER SEQ ID NO: 24

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 358 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

5

10

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

15

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

20

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

25

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

30

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 24

```

aatgggggac cggagaagaa gtacaagagc acgggtcagg ggcggggactc cgacgggetcc 60
ttcttctctt acagcaggct aaccgtggac aagagcagggt ggcaggagggt gaatgtcttc120
tcatgtctcg tgatgcatga ggctctgcac aaccactaca cgcagaagag cctctccctg180
tctccgggta aatgagtgcg acggccggca agcccccgct ccccgggctc tcgcggtcgc240
acgaggatgc ttggcacgta ccccggtgtac atacttccca ggcacccagc atggaaataa300
agcaccacgc gctgccctgg ggccccctgcg aaaaaaaaga aaaagaatcg aaaagggg 358

```

40

(2) INFORMATION ÜBER SEQ ID NO: 25:

45

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 89 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

50

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

55

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

60

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

65

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 25

gcccctagcc cctggcagac atagctgctt cagtgtcccct tttcctctgc tggctagatg60
 gatgttgatg cactggaggt acttttagc 89

(2) INFORMATION ÜBER SEQ ID NO: 26:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1632 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 26

gacactgggtt ggttctgata agaggcaggg gaggagaaag ccgaggaaga gggagttgcg 60
 gaagaggagg gagttaacaa gttctcttat ccaccatcac accgggagtg ttgtccagcc 120
 gtggaggagg aggacgatga agaagctgta aagaaagaag ctacacagaac ctctacttct 180
 45 gcttgtctc caggatccaa gccagcact tgggtgtctt gccagggga ggaagagaat 240
 caagccacgg aggataaaaag aacagaaaga agtaaaaggag ccaggaagac ctccgtgtcc 300
 ccccgatctt caggctccga ccccggtcc tgggagtatc gttcaggaga ggcgtccgag 360
 gagaaggagg aaaaggcaca caaagaaact gggaaaggag aagctgcccc agggccgcaa 420
 50 tcttcagccc cagcccagag gccccagctc aagtcctggt ggtgccaacc cagtgatgaa 480
 gaggagggtg aggtcaaggc tttgggggca gctgagaagg atggagaagc tgagtgtcct 540
 ccctgcatcc cccaccagag tgccttcttg aaggcctggg tgtattggcc aggagaggac 600
 acagaggaag aggaagatga ggaagaagat gaggacagtg actctggatc agatgaggaa 660
 gaggagaaag ctgaggcttc ctcttcact cctgctacag gtgtcttctt gaagtcctgg 720
 55 gtctatcagc caggagagga cacacagtga tacaggatca gccgaggatg aaagagaagc 780
 tgagacttct gcttccacac cccctgcaag tgctttcttg aaggcctggg tgtatcggcc 840
 aggagaggac actggatagt gaggataagg aagatgattc agaagcagcc ttaggagaag 900
 ctgagtcaga cccacatccc tcccaccgg accagagggc ccacttcagg ggctggggat 960
 atcgacctgg aaaagagaca gaggaagagg aagctgctga ggactgggga gaagctgagc1020
 60 cctgccccct cggactggcc atctatgtac ctggagagaa gccaccgctt ccttgggctc1080
 cctctaggct gccctccga ctgcaaaggg ggtcaagcg ccagaaacc cctactcatg1140
 atccggaccc tgagactccc ctaaaggcca gaaagggtgcg cttctccgag aaggtcactg1200
 tccatttctt ggctgtctgg gcagggccgg cccaggccgc ccgccagggc ccctgggagc1260
 agcttgctcg ggatgcgagc cgcttcgcac gccgcatcac ccaggcccag gaggagctga1320
 65 gccctgcct caccctgct gcccgggcca gacgtgggc acgctcagg aaccacctt1380
 tagcccccct cctgcccctc acccagacct tgccttctc ctctgtccct tcgtccccag1440
 tccagaccac gcccttgagc caagctgtgg ccacacctc ccgctcgtct gctgctgcag1500
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ttactatttt attttttcta agtgtgggtt tatataagga ataaagcctt ttgatttgta1620
acgaaaaaaa aa 1632

(2) INFORMATION ÜBER SEQ ID NO: 27:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 2972 Basenpaare
(B) TYP: Nukleinsäure
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library.

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 27

ccaggacgag cacctcatta cattcttcgt gcctgtcttt gagccgctgc cccctcagta 60
cttcatccga gtggtgtctg accgctggct ctcttgtgag acccagctgc ctgtctcctt 120
ccggcacctg atcttgccgg agaagtaccc cctcccaacc gaacttttgg acctgtagcc 180
cttgcccgtg tctgtctga gaaacagtgc ctttgagagt ctttaccag ataaatttcc 240
tttcttcaat cccatccaga cccaggtgtt taacactgta tacaacagt acgacaacgt 300
gtttgtgggg gccccacgg gcagcgggaa gactatttgt gcagagtttg ccatcctgcg 360
aatgctgctg cagagctcgg aggggcgctg tgtgtacat acccccatgg aggcctggc 420
agagcagga tacatggact ggtacgagaa gttccaggac aggtcaaca agaaggtggt 480
actcctgaca ggcgagacca gcacagacct gaagctgctg ggcaaaggga acattatcat 540
cagcacccct gagaagtggg acatacttcc ccggcgatgg aagcagcgca agaactgca 600
gaacatcaac ctcttcgtgg tggatgaggt ccacctatc gggggcgaga atgggcctgt 660
cttagaagt atctgtctcc gaatgcgcta catctctcc cagattgagc ggccattcg 720
cattgtggca ctcagctctt cgctctccaa tgccaaggat gtggccact ggctgggctg 780
cagtgccacc tccaccttca acttccatcc caatgtgct cccgtccctc tggagctgca 840
catccagggc ttcaacatca gccatacaca aaccgcctg ctctccatgg ccaagcctgt 900
gtaccatgct atcaccaagc actcgcccaa gaagcctgtc attgtctttg tgcctctcg 960
caagcagacc cgcctcactg ccattgacat cctcaccacc tgtgcagcag acatccaacg1020
gcagaggttc ttgcaactga ccgagaagga tctgattccg tacctggaga agctaagtga1080
cagcacgctc aaggaaacgc tgctaaatgg ggtgggctac ctgcatgagg ggctcagccc1140
catggagcga cgcctgggtg agcagctctt cagctcaggg gctatccagg tgggtggggc1200
ttctcggagt ctctgctggg gcatgaacgt ggctgcccac ctggtaataca tcatggatac1260
ccagtactac aatggcaaga tccacgccta tgtggattac cccatctatg acgtgcttca1320
gatgggtggc cagccaacc gccctttgca ggacgatgag gggcgctgtg tcatcatgtg1380
tcagggtccc aagaaggatt tcttcaagaa gttcttatat gagccattgc cagtagaatc1440
tcacctggac cactgtatgc atgaccactt caatgctgag atcgtcacca agaccattga1500
gaacaagcag gatgctgtgg actacctcac ctggaccttt ctgtaccgcc gcatgacaca1560
gaaccceaat tactacaacc tgcagggcac ctcccatcgt cacttgctgg accactgtgc1620
agagctgggt gagcagaccc tgaagtacct ggagcagtc aagtgcata gcatcgagga1680
cgagatggac gtggcgctc tgaacctagg catgatcgcc gcctactatt acatcaacta1740
caccaccatt gagctcttca gcatgtccct caatgccaa accaaggtgc gagggcttat1800

cgagatcatc tccaatgcag cagagtatga gaacatccc atccggcasc atgaagacaa1860
 tctcctgagg cagttggctc agaaggtccc ccacazgctg aataacccta agltcaatga1920
 tccgcacgtc aagaccaacc tgctcctgca ggctcacttg tctcgcatgc agctgagtgc1980
 5 tgagttgcag tcagatacgg aggaatcct tagtaaggca atccggctca tccaggcctg2040
 cgtggatgtc ctttccagca atgggtggct cagccctgct ctggcagcta tggaaactggc2100
 ccagatggtc acccaagcca tgtggtccaa ggactcatac ctgaagcagc tgccacactt2160
 cacctctgag catatcaaac gttgcacaga caaggagtg gagagtgtt tcgacatcat2220
 10 ggagatggag gatgaagaac ggaacgcgtt gcttcagctg actgacagcc agattgcaga2280
 tgtggctcgc ttttgtaacc gctaccctaa tatcgaacta tcttatgagg tggtagataa2340
 ggacagcatc cgcagtggcg ggccagtgtg ggtgctggtg cagctggagc gagaggagga2400
 agtcacaggc cctgtcattg cgctctctt cccgcagaaa cgtgaagagg gctgggtggg2460
 ggtgattgga gatgccaagt ccaatagcct catctccatc aagaggctga ccttgacagca2520
 15 gaaggccaag gtgaagttgg actttgtggc ccagccact ggtgcccaca actacactct2580
 gtacttcatg agtgacgctt acatgggatg tgaccaggag tacaattca gcgtggatgt2640
 gaaagaagct gagacagaca gtgattcaga ttgagtcctg aggcatttac ttttgggtaa2700
 aggagagttg agcctgaatt aggaatgtgt acattgtagg aatcctggtt gtggggacca2760
 ggtctgtggg cctcaggtct ggccagccag ggctggtgct gtccccgcct acctccactt2820
 20 cctttccctt gctcactctg gatccagtga cagcaggtgt catgggtcaa gcataaatca2880
 tatatagcat tttcaggcat gttcctggta gttcttttga gtctgacatt ctaataaaat2940
 aatttgtaga aaaaaaacca aaaaaaaaaa aa 2972

25 (2) INFORMATION ÜBER SEQ ID NO: 28:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 496 Basenpaare
 30 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

35 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

40 (iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- 45 (A) ORGANISMUS: MENSCH
 (C) ORGAN:

50 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 28

55 ctcgagccga agagtccttg acctcctgtg caagaacatg aaacatctgt ggttcttctt 60
 tctcctgggtg gcagctccca gatgggtcct gtcccagggt cagctgcagg agtcggggccc120
 aggactgggtg aagccttcgg agaccctgtc cctcacctgc actgtctctg gtggctccat180
 60 cagtagttac tactggagct ggatccggca gccccaggg aagggactgg agtggattgg240
 gtatatctat tacagtggga gcaccaacta caaccctcc ctcaagagtc gagtacccat300
 atcagtagac acgtccaaga accagttctc cctgaagctg agctctgtga ccgctgcgga360
 cacggccgtg tattactgtg cgagacaggg tatagcagtg gaccagcttg actactgggg420
 ccagggaacc ctggtcaccg tctcctgagc ctgcaccaag gggccatcgg tcttccccct480
 65 ggcaccctgc tccaag 496

(2) INFORMATION ÜBER SEQ ID NO: 29:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 397 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

10

(iii) HYPOTHETISCH: NEIN

15

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

20

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

25

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 29

gaggctcctgg acctcctgtg caagaacatg aaacacctgt ggttcttcct cctcctgggtg 60
 gcagctccca gatgggtcct gtcccagggtg cagctgcagg agtcggggccc aggactgggtg120
 aagccttcgg agaccctgtc cctcacctgc actgtctctg gtggctccat cagtagttac180
 tactggagct ggatccggca gccgcgggg aagggactgg agtggattgg gcgtatctat240
 accagtggga gcaccaacta caaccctcc ctcaagagtc gagtcaccat gtcagtagac300
 acgtccaaga accagttctc cctgaagctg agctctgtga ccgccgcgga cacggccgtg360
 tattactgtg cgagagcaaa acgcagctgg acctcag 397

30

35

(2) INFORMATION ÜBER SEQ ID NO: 30:

40

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 772 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

45

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

50

(iii) HYPOTHETISCH: NEIN

55

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

60

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

65

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 30

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5  gggactcctc agttcacctt ctcacaatga ggctccctgc tcagtcctg gggctgctaa 60
   tgctctgggt ctctggatcc agtggggata ttgtgatgac tcagtcctca ctctccctgc 120
   ccgtcaccctc tggagagccg gcctccatct cctgcaggtc tagtcagagc ctctgcata 180
   gtaatggata caactatttg gattggtacc tgcagaagcc agggcagctc ccacagctcc 240
   tgatctatctt ggggtctaat cgggcctccg gggtcctga caggttcagt ggcagtgat 300
10  caggcacaga ttttactactg aaaatcagca gagtggaggc tgaggatgtt ggggtttatt 360
   actgcatgca agctctacaa actcctctca ctttcggcgg agggaccaag gtggagatca 420
   aacgaactgt ggctgcacca tctgtcttca tcttcccgcc atctgatgag cagttgaaat 480
   ctggaactgc ctctgttggtg tgctgtctga ataacttcta tccagagag gccaaagtac 540
15  agtggaaggt ggataacgcc ctccaatcgg gtaactccca ggagagtgtc acagagcagg 600
   acagcaagga cagcacctac agcctcagca gcaccctgac gctgagcaaa gcagactacg 660
   agaaacacaa agtctacgcc tgcgaagtca cccatcaggg cctgagcttg gcccgtaa 720
   aaagggtctt caacaggggg aagtttttag aggggagatg tggccccacc tt 772

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(2) INFORMATION ÜBER SEQ ID NO: 31:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 1031 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 31

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55  ggggacggaa gcgcagagca cggaccccg cccctcgcg ccccgctcgt gacgtcgcg 60
   ggggcccgg cctccgccg gccccgagg gctctcccg gaggtcagc cccctctgt 120
   ccccatgggc aactgccagg cagggcacaa cctgcaactg tgtctggccc accaccacc 180
   tctggtctgt gccactttga tctgtctgt ccttggtctc tctggcctgg gccttggcag 240
   ctctcctctc acccacagga ctggcctgcg cagcctgaca tccccagga ctgggtctct 300
   tttttgagat cttttggcca gctgaccctg tgtcccagga atgggacagt cacagggag 360
60  tggcgagggt ctcacgtcgt gggcttgcgt accaccttga acttcggaga cggtcagac 420
   aggaacaaga cccggacatt ccaggccaca gtccctggga gtcagatggg attgaaagga 480
   tcttctgcag gacaactgtt ccttatcaca gccagggtga ccacagaaag gactgcagga 540
   acctgcctat attttagtgc tgttccagga atcctacct ccagccagcc acccatatcc 600
   tgctcagagg agggggctgg aaatgccacc ctgagcccta gaatgggtga ggaatgtgtt 660
65  agtgtctgga gccatgaagg ccttgtgtct accaagctgc tcacctcga ggagctggct 720
   ctgtgtggct ccaggctgct ggtcttgggc tcttctctgc ttctctctc tggccttctc 780
   tgctgtgtca ctgctatgtg cttccaccgc cgccgggagt cccactggtc tagaaccgg 840
   ctctgagggc actggcctag ttcccgaact gtttctcagg tgtgaatcaa cttcttgggc 900

```

cttggtctctg agttggaaaa ggttttagaa aaactgaaga gctggatgt ggaggaaaa 960
 aaaagcttt tttgcccaa aaaaaaaaaa aaaaaaa aaaaaaa aaaaaaa 1020
 aaaaaaaaa a 1031

5

(2) INFORMATION ÜBER SEQ ID NO: 32:

(i) SEQUENZ CHARAKTERISTIK:

10

(A) LÄNGE: 739 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

15

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

20

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

25

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

30

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

35

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 32

cggtctgagc cccgtctcagt caccgcgagc aggcgtgcag tttcccggt ctccgcgcgg 60
 ccggggaagg tcagcccggt aatggcggtc ttggcgctcg gacctacct gacctatcag120
 caaaaggtgt tgcggcttta taagcgggag ctacgccacc tcgagtcgtg gtgcgtccag180
 agagacaaat accgatactt tgcttggttg atgagagccc ggtttgaaga acataagaat240
 gaaaaggata tggcgaaggc caccagctg ctgaaggagg ccgaggaaga attctggtac300
 cgctcagcatc cacagccata catcttcctt gactctcctg ggggcacctc ctatgagaga360
 tacgattgct acaaggtccc agaattgtgc ttagatgact ggcacacctc tgagaaggca420
 atgtatcctg attactttgc caagagagaa cagtgggaaga aactgcggag ggaaagctgg480
 gaacgagagg ttaagcagct gcaggaggaa acgccacctg gtggtccttt aactgaagct540
 ttgccccctg cccgaaagga aggtgatttg cccccactgt ggtggtatat tgtgaccaga600
 ccccgggagc ggcccatgta gaaagagaga gacctcatct ttcatgcttg caagtgaat660
 atgttacaga acatgcactt gccctaataa aaaatcagtg aaatggaaaa aaaaaaaaaa720
 aaaaaaaaa aaaaaaaaa 739

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(2) INFORMATION ÜBER SEQ ID NO: 33:

(i) SEQUENZ CHARAKTERISTIK:

60

(A) LÄNGE: 651 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

65

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 33

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cggtctgagc ctctagttcac cttctcacca tgaggctccc tgctcagctc ctggggctgc 60
taatgctctg ggtccctgga tccagtggag atattgtgat gacctcagact ccactctccc120
tgcccgtcac ccctggagag ccggcctcca tctcctgcag gtctagtcag agcctcttgg180
atagtgatga tggaaacacc tatctggact ggtacctgca gaagccaggg cagtctccac240
agctcctgat ctatacgctt tcctatcggg cctctggagt ccagacaggg ttcagtggca300
gtgggtcagg cactgatttc aactgaaaa tcagcagggg ggaggctgag gatgttggag360
tttattactg catgcaacgt atagaatttc cttacacttt tggccagggg accaagctgg420
agatcaaacg aactgtggct gcaccatctg tcttcactct cccgccatct ggatgagcag480
ttgaaatctg gaacttgccct ctgttgttgt gcctgcttga ataactttct attcccagag540
aggggcaaag taacagtgga aggttggatt aacgccctgc aattcgggta actgccagg600
gagtagtttt cacagggcag gggcagcaag gacagcacct acagtcttag t 651

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(2) INFORMATION ÜBER SEQ ID NO: 34:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 823 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 34

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ccgcgtcgac aaattttttt aaagatcatc gatgaagaga gaaaatgcgc ttttctacag 60
agtcccttc ccaccacag ccccatcccc agataagcgg ggagttccct ggcgcggtgc120
cagtttctag ccgctgagtg ggcgtgtgcg cggctccaag tgcgcctgcg tactgctcac180
tccccagctc cgcgccttgc tccgttcctc ccaaaactct gaatcgaaga actttccgga240
agtttctgag agcccagacc ggcgggcaag cgcccatccc caacccctc tgttaatccc300
taccagcctg cagtccctggc tgcttccaag caggaggtgg ggcctctggc ctagcggggc360

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cgaaaggcag tgeccctccc ccgcagtctg atttccctct tccccccaac ggcaagcacg120
aggagcggca ggacgagcat ggctacatct cccggtgctt cacgcggaac tacacgtgc480
cccccggtgt ggaccccacc caagtttctt cctccctgtc ccctgagggc acactgaccg540
tgagggcccc catgcccaag cttagccacgc agtccaacga gatcaccatc ccagtcacct600
tcgagtcgcg ggcccagctt gggggcccag aagctgcaaa atccgatgag actgccgcca660
agtaaagcct tagcccgat gcccccctt gctgccgcca ctggctgtgc ctcccccgcc720
acctgtgtgt tcttttgata cttttatctt ctgtttttct caaataaagt tcaaagcaac780
cacctggtca aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa aaa 823

5

10

(2) INFORMATION ÜBER SEQ ID NO: 35:

(i) SEQUENZ CHARAKTERISTIK:

15

(A) LÄNGE: 457 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

20

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

25

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

30

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

35

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

40

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 35

cataagggaa atgctttctg agagtcacatg atctcatgtg caagaaaatg aagcacctgt 60
ggttcttctt cctgctgggt gcggtcccca gatgggtcct gtcccagctg cagctgcagg120
agtcggggccc aggactgggt aagccttcgg agaccctgtc cctcacctgc actgtctctg180
gtggctccat cagcagtagt agttactact ggggctggat ccgccagccc ccagggaagg240
ggctggagtg gattgggagt atctattata gtgggagcac ctactacaac ccgtccctca300
agagtcgagt caccatatcc gtagacacgt ccaagaagta cttctccctg aagctgagct360
ctgtgaccgc cgcagacacg gctgtgtatt actgtgcgag acatgactgg tattacgata420
ttttgactgg ttatgcgaaa cccggcacag gtccgac 457

45

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(2) INFORMATION ÜBER SEQ ID NO: 36:

55

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 1203 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

60

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

65

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 36

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gtcggggg cctgcgcagt cgctcttctt caggcggcgg ccatggcggg acaggaggat 60
ccggtgcagc gggagattca ccaggactgg gctaaccggg agtacattga gataatcacc 120
agcagcatca agaaaatcgc agactttctc aactcgttcg atatgtcttg tcgttcaaga 180
cttgcaacac taaacgagaa attgacagcc cttgaacgga gaatagagta cattgaagct 240
cgggtgacaa aagggtgagac actcacctag aacagtgccg tgcgtctgct ggggaagttgc 300
tttacacaac acaggccaca tgggaaaggc cccagcagcc ttcagctcct tcctttctcc 360
ttaaagagca acagggctta ttcttggttt tcttttttca aaagtgtggc ctttgggctc 420
tgccatctgg ggtgtggtgt ggtatgtggg aagaagttca gaggaaccgt tggaaacgac 480
gttaggcatt ttaccttttc agtaacattt tatacatcta cttgtcaatg tatttgagac 540
attcacagcc aaaagcctgg gactctttgt gaaggctctc ctcacctcta tctttctttc 600
tctctctctc aaactttcct taaagttctc attgcctttg cactgcttct gtgaacagtc 660
tttgtctcct cccacctttt ggtgggaagt gcggggcagt cctgggtcaag acactcatgc 720
cctggcaatg tggctgccag agaattgttg tgctaaccga ccagtttctt gttgatttgg 780
agagggtcaag gccaggcccc cacttggttt gaagggacat tttcagactt ttctttctgt 840
cacttgaggt gtctatgcct ctcataattc cctaataaac tcctcaactt tttatctgac 900
tgctgtgatt atggtgggga gaggagctag agatgggttc acttattgca cagaaatgta 960
atacatggcg ttattattct aacataaaac tttcagatgt agctgtttga ttcaaagcct1020
aggtgcttac cagcccaagt ccccatgttt ggactttcag ctgactagct catcttggga1080
atcatttggg cattcagcac atttaccag tatttactat gtaggcattg taaactcca1140
taaacataac agcattgaat cagaaaaaaa aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa1200
aaa

```

(2) INFORMATION ÜBER SEQ ID NO: 37:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 207 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 37

cggctcgagg cgcgcctcgg tgtcagtgtc cccaggacag acggccagga tcacctgtc 60
 tggacatgca ttgccaaagc aatgatgctt attggtacca gtcagaggcc agggccaggc120
 cctgtgctt ggtggatccc ttgaaagaac attggaggag ggccttcag ggcatgccct180
 ggagacgat tgctctgggc ttccaac 207

(2) INFORMATION ÜBER SEQ ID NO: 38:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 346 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 38

cggctcgagc ggctcgaggc cgggatggtg ggtgctacgc cccttgggta ctggggccag 60
 ggaaccctgg tcaccgtctc ctcagcctcc accaagggcc catcgggctt cccccgggca120
 cctcctcca agagcacctc tgggggcaca gcggcctgg gctgcctggt caaggactac180
 ttccccgaac cggtagcggg gtcgtggaac tcaggcgctt gaccagcggc gtgcacacct240
 tccccgctgt ctacagtctc aggactctac tcctcagcag cgtggtgacg tgccctccag300
 cagttgggca ccagacctac atctgcaagt gaatcgaagc cagcaa 346

(2) INFORMATION ÜBER SEQ ID NO: 39:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 926 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 39

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cggtcttaag gaagcagcac tgggtggtgcc tcagccatgg cctggaccgt tctcctcctc 60
ggcctcctct ctactgcac aggtctctgtg acctcctatg tgctgactca gccaccctcg120
gtgtcagtg cccacaggaca gacggccagg attacctgtg ggggaaacaa cattggaagt180
aaaagtgtgc actggtacca gcagaagcca ggccaggccc ctgtgctggt cgtctatgat240
gatagcgacc ggccctcagg gatccctgag cgattctctg gctccaactc tgggaacacg300
gccaccctga ccatcagcag ggtcgaagcc ggggatgagg ccgactatta ctgtcaggtg360
tgggtagtga gtagtgatca ttgggtgttc ggcggaggga ccaagctgac cgtcctagg420
cagcccaagg ctgccccctc ggtcactctg ttcccgcct cctctgagga gcttcaagcc480
aacaaggcca cactggtgtg tctcataagt gacttctacc cgggagccgt gacagtggcc540
tgggaaggcag atagcagccc cgtcaaggcg ggagtggaga ccaccacacc ctccaaacaa600
agcaacaaca agtacgcggc cagcagctat ctgagcctga cgcctgagca gtggaagtcc660
cacagaagct acagctgcca ggtcacgcat gaaggagca ccgtggagaa gacagtggcc720
cctacagaat gttcataggt tctcaaccct cccccccac caccgggagac tagagtgc840
ggatcccagg ggagggtct ctctccac cccaaggcat caagccctc tccctgcact840
caataaacc tcaataaata ttctcattgt caatcaggaa aaaaaaaaaa aaaaaaaaaa900
aaaaaaaaaa aaaaaaaaaa aaaaaa

```

926

(2) INFORMATION ÜBER SEQ ID NO: 40:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 2384 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 40

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gcctcccgcc cgcgcctct gtctccctct ctccacaaac tgcccaggag tgagtagctg 60
ctttcggtcc gccggacaca ccggacagat agacgtgcgg acggcccacc accccagccc 120
gccaactagt cagcctgccc ctggcgctc ccctctccag gtccatccgc catgtggccc 180

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ctgtggcgcc	tcgtgtctct	gttggccctg	agccaggccc	tgccttttga	gagagagagge	210
ttctgggact	tcaccttgga	cgatggggcca	ttcatgatga	acgatigagga	agcttcgggc	300
gctgacacct	cgggcgctct	ggaccgggac	tctgtcacac	ccacctacag	cgccatgtgt	360
cctttcgggt	gccactgcca	cctgcggtgt	gttcagtgtc	ccgacctggg	tctgaagtct	420
gtgcccagaag	agatctcccc	tgacaccacg	ctgctggacc	tgagaacaa	cgacatctcc	480
gagctccgca	aggatgactt	caagggtctc	cagcacctct	acgccctcgt	cctgggtgaac	540
aacaagatct	ccaagatcca	tgagaaggcc	ttcagcccac	tgcggaagct	gcagaagctc	600
tacatctcca	agaaccacct	ggtggagatc	ccgcccaccc	taccagctc	cctgggtggag	660
ctccgcatcc	acgacaaccg	catccgcaag	gtgcccgaag	gagtgttcag	tggtgctccg	720
aacatgaact	gcatcgagat	ggcggggaac	ccactggaga	acagtggcct	tgaacctgga	780
gccttcgatg	gcctgaagct	caactacctg	cgcatctcag	aggccaagct	gactggcatc	840
cccaaagacc	tccttgagac	cctgaatgaa	ctccacctag	accacaacaa	aatccaggcc	900
atcgaactgg	aggactgtct	tcgctactcc	aagctgtaca	ggctgggcct	aggccacaac	960
cagatcagga	tgatcgagaa	cgggagcctg	agcttcctgc	ccacctccg	ggagctccac	1020
ttggacaaca	acaagttggc	cagggtgccc	tcagggtctc	cagacctcaa	gctcctccag	1080
gtggtctatc	tgactccaa	caacatcacc	aaagtgggtg	tcaacgactt	ctgtcccatg	1140
ggcttcgggg	tgaagcgggc	ctactacaac	ggcatcagcc	tcttcaacaa	ccccgtgcccl	1200
tactgggagg	tgacgcccgc	cactttccgc	tgcgctcactg	actgcctggc	catccagttt	1260
ggcaactaca	aaaagtagag	gcagctgcag	ccaccgcggg	gcctcagtgg	gggtctctgg	1320
ggaacacagc	cagacatcct	gatggggagg	cagagccagg	aagctaagcc	agggccagc	1380
tgcgtccaac	ccagccccc	acctcgggtc	cctgacccca	gctcgatgcc	ccatcacgc	1440
ctctccctgg	ctcccaagg	tgagggtggg	cgcaaggccc	ggccccatc	acatgttccc	1500
ttggcctcag	agctgcccct	gctctccac	cacagccacc	cagaggcacc	ccatgaagct	1560
tttttctcgt	tactcccaa	acccaagtgt	ccaaggctcc	agtcctagga	gaacagtccc	1620
tggtgcagca	gccaggaggc	ggtccataag	aatggggaca	gtgggtctctg	ccagggtcgc	1680
cgacacctgtc	cagacacaca	tgttctgttc	ctcctcctca	tgcatttcca	gcctttcaac	1740
cctccccgac	tctgcggtc	ccctcagccc	ccttgcaagt	tcatggcctg	tcctcccag	1800
accctgtctc	cactggccct	tcgaccagtc	ctccctctctg	ttctctcttt	ccccgtcctt	1860
cctctctctc	tctctctctc	tttctgtgtg	tgtgtgtgtg	tgtgtgtgtg	tgtgtgtgtg	1920
tgtgtgtgtc	ttgtgtctcc	tcagaccttt	ctcgcttctg	agcttgggtg	cctgttccct	1980
ccatctctcc	gaacctggct	tcgcctgtcc	ctttcactcc	acacctctg	gccttctgcc	2040
ttgagctggg	actgctttct	gtctgtccgg	cctgcaccca	gccccgtccc	acaaaacccc	2100
agggacagcg	gtctccccag	cctgccctgc	tcaggccttg	cccccaaacc	tgtactgtcc	2160
cggaggagggt	tgggagggtg	aggcccgaca	tcccgcgag	atgacacccat	caaccgcccag	2220
agtcccagac	accggttttc	ctagaagccc	ctcaccacca	ctggcccaact	ggtggctagg	2280
tctcccctta	tcttctctgt	ccagcgcaag	gaggggctgc	ttctgagggtc	ggtggctgtc	2340
ttccattaa	agaaacaccg	tgcaacgtga	aaaaaaaaa	aaaa		2384

(2) INFORMATION ÜBER SEQ ID NO: 41:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 334 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

5 (xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 41

ctcgagccga attcggtctg agaggagccc agccctggga ttttcaggtg ttttcatttg 60
 gtgatcagga ctgaacagag agaactcacc atggagtttg ggctgagctg gctttttctt120
 10 gtggctattt taaaagggtg ccagtgtgag gtgcagctgt tggagtctgg gggaggcttg180
 gtacagcctg gggggtcctt gagactctcc tgtgcagcct ctggattcac cttagcagc240
 tatgccatga gctgggtccg ccagggtcca gggaaggggc tggagtgggt ctcaggtatt300
 agtggtagtgt gtgtgatagt acacactacg caga 334

15 (2) INFORMATION ÜBER SEQ ID NO: 42:

(i) SEQUENZ CHARAKTERISTIK:

- 20 (A) LÄNGE: 845 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

25 (ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

30 (iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

35 (vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

40 (vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

45 (xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 42

gcgttccctc cgccgagcta cttctttctt tccttttttt tttttttctg gctaacagaa 60
 ttttattggt aaatcacaga aacttttagt caaaacaaaa atcacgaagt ccattttaata120
 gcaacttcat gtctgtctgg ctttgcttgc tgtctcctgg caaccagaag tggacagaag180
 50 cgtgggtgcc caagtgggcc acagacagct tccaaccccc acacccagc atccaatcca240
 caccagcag acccttcgcc atgccgccct ctaccaggaa gccagaggcc taggagctcg300
 ccattccatat ttatttgaaa aggtcaaaag gaggcatctat gagacaaggg aggggtgcag360
 gctgaagcag cgctcaaca gccagggaca ttaggcaac acgagcaggc acagcgcggc420
 caccactgtc cacacgctca cacaagccag gcccgaggc ccttcggaga gctagcaggt480
 55 tacattcagg cagatggccc tcttcccacc caaaccaca gaaccccaa caaggcatca540
 ccaggaaaga cacgggaaag ccaaatcaca gttgaaccag ggacagagaa cccttggccc600
 cactgatgtc ccaagccacc agcagctgtc tccaaaatcc ctatgctatt acagtgggaa660
 ttacatcatt taaaaagcct gattattccc aggettctaa tctttcatat aaaactgcct720
 60 ttgttttgct cctttgttca actcagaggc ccagcaaaagc gggcagggtc cctgatcagg780
 gcaggagccc acctcagaag cccatgccgc accagtgcgc aagcacatgt cagtgtcag840
 aacaa 845

65 (2) INFORMATION ÜBER SEQ ID NO: 43:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 2233 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

10

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

15

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH
 (C) ORGAN:

20

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

25

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 43

gaattcagaa	gttaaatgatg	ttgggtaaga	gaacaatggt	aagagagcaa	tctaagaata	60
tatcacctac	tttaatttta	tatgagagta	catggaggta	gctgtgatgt	ggaaatgtag	120
cactgctcct	acccacgcag	atttattcca	gtgaaacaac	aactggaact	tcaagtaact	180
cctcccagag	tacttccaac	tctgggttgg	ccccaatcc	aactaatgcc	accaccaagg	240
cggctggtgg	tgccctgcag	tcaacagcca	gtctcttcgt	ggctctcactc	tctcttctgc	300
atctctactc	ttaagagact	caggccaaga	aacgtcttct	aaatttcccc	atcttctaaa	360
cccaatccaa	atggcgctcg	gaagtccaat	gtggcaagga	aaaacaggtc	ttcatcgaat	420
ctactaattc	cacacctttt	attgacacag	aaaatggtga	gaatcccaaa	tttgattgat	480
ttgaagaaca	tgtgagaggt	ttgactagat	gatggatgcc	aatattaaat	ctgctggagt	540
ttcatgtaca	agatgaagga	gaggcaacat	ccaaaatagt	taagacatga	tttccttgaa	600
tgtggcttga	gaaatattga	cacttaatac	taccttgaaa	ataagaatag	aaataaagga	660
tgggattgtg	gaatggagat	tcagttttca	tttggttcct	taattctata	aggccataaa	720
acaggtaata	taaaaagctt	ccatgattct	atztatatgt	acatgagaag	gaacttccag	780
gtgttactgt	aattcctcaa	cgtattgttt	cgacagcact	aatttaatgc	cgatatactc	840
tagatgaagt	tttacattgt	tgagctattg	ctgttctctt	gggaactgaa	ctcactttcc	900
tectgaggct	ttggatttga	cattgcattt	gaccttttat	gtagtaattg	acatgtgcca	960
gggcaatgat	gaatgagaat	ctacccccag	atccaagcat	cctgagcaac	tcttgattat	1020
ccatattgag	tcaaattgta	ggcatttcct	atcacctggt	tccattcaac	aagagcacta	1080
cattcattta	gctaaacgga	ttccaaagag	tagaattgca	ttgaccgcga	ctaatttcaa	1140
aatgcttttt	attattatta	tttttttagac	agtctcactt	tgtcgcccag	gccggagtgc	1200
agtgtgcgca	tctcagatca	gtgtaccatt	tgcctcccgg	gctcaagcga	ttctcctgcc	1260
tcagcctccc	aagtagctgg	gattacaggc	acctgccacc	atgcccggt	aatttttgta	1320
attttagtag	agacagggtt	tcaccatgtt	gccaggctg	gtttcgaact	cctgacctca	1380
ggtgatccac	ccgcctcggc	ctoccaaagt	gctgggatta	caggcttgag	cccccgcgcc	1440
cagccatcaa	aatgcttttt	atctctgcat	atggtgaata	ctttttacaa	tttaaaaaaa	1500
tgatctgttt	tgaaggcaaa	attgcaaatc	ttgaaattaa	gaaggcaaaa	atgtaaagga	1560
gtcaaaacta	taaatcaagt	atttgggaag	tgaagactgg	aagctaattt	gcattaaatt	1620
cacaaacttt	tatactcttt	ctgtatatac	attttttttc	tttaaaaaac	aactatggat	1680
cagaatagcc	acattttaga	cactttttgt	tatcagtcac	taatttttaga	tagttagaa	1740
ctggtcctaa	gcctaaaagt	gggcttgatt	ctgcagtaaa	tctttttacaa	ctgcctcgac	1800
acacataaac	cttttttaaaa	atagacactc	cccgaagtct	tttgttcgca	tggtcacaca	1860
ctgatgctta	gatgttccag	taatctaata	tggccacagt	agtcttgatg	accaaagtcc	1920
tttttttcca	tcttttagaaa	actacatggg	aacaaacaga	tcgaacagtt	ttgaagctac	1980
tgtgtgtgtg	aatgaacact	cttgcttttat	tccagaatgc	tgtacatcta	ttttggattg	2040
tatatgtgtg	ttgtgtattt	acgctttgat	tcatagtaac	ttcttatgga	attgatttgc	2100
attgaacaca	aactgtaaat	aaaaagaaat	ggctgaaaga	gcaaaaaaaa	aggagggcag	2160
gagagaggaa	aaggggagga	agaggagggg	ggaaagagaa	gggagagaga	aggaggggga	2220
aggaggtggg	ggg					2233

65

(2) INFORMATION ÜBER SEQ ID NO: 44:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 243 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 44

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ggagcccagc actagaagtc ggcgggtgttt ccattcggtg atcagcactg aacacagagg 60
actcaccatg gagtttgggc tgagctgggt tttcctcgtt gctcttttaa gaggtgtcca120
gtgtcagggtg cactggtgga gcggggagcg ggtcagcagg agtcctgaat cctgtgacgc180
tgatcagtc tatatcagat ggcgcagctc agcagggtga tggggtatga atgataacat240
aca                                     243

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(2) INFORMATION ÜBER SEQ ID NO: 45:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 817 Basenpaare
- (B) TYP: Nukleinsäure
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
- (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 45

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gttttttttt tttttttttg aagagagcag attctcttta ttgagatacg ggacacagcg 60
aagggtggag agacggaaca gccccccagc ctcagccctc tccacggggg ccggtatgcca120
gagatgggag aagggtattca gtctctcgcc cgggaaacce agtcccacag agggcgccgg180
caagggtggg acgcgacctg ggtgacacgg tgcaggaggt ctttaaatag aggaggggct240
ggagcgggga aacgcgccgg gggcctagcg caccatgtat tccttgcgct tattgagccg300
aacttggcag aaagagaagc ctccgaggag gaggtaaagg cctgcagcga tgaaacagtt360
gtagctgact tgctcgtaaa ggttgatat gttctggggg ccattctcaa aatctttctc420
cgtgaaggga acgtcctcaa tcaacacagc ggaatggaca ttgaaaaata ttccgagcat480
tatcaacatg atcactcccc aggcgctgag gacgatgccg caggcggccca gcttcggccc540
acagcacagg agcgacgcca taaagaagg agtcggggat cgccgagggt caagcgggct600
cggaagcgg tgggagaaag cccaggatgc cctcgagggg ggcagaggg ggcgtggccc660
cgccctcaac catcccatcc gggggcgcca ggcggaaaag gctgggctcc tctcaggact720
ttcgcgggag acggcgccgt ctgaaaccaa aactgtcctt ggggaaacct tccttgacct780
ctgtagctag ggcgtgagta ttggaagagc gagggccc 817

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(2) INFORMATION ÜBER SEQ ID NO: 46:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 1644 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 46

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gttccggctc acatgggaaa tactttctga gagtcctgga cctcctgtgc aagaacatga 60
aacacctgtg gttcttcctc ctgctgggtg cagctcccag atgggtcctg tcccagggtgc 120
agctgcagga gtcggggcca ggactgggtg agccttcaca gacctgtcc ctcacctgca 180
ctgtctctgg tggtccatc agcagtgggt gttactactg gagctggatc cgccagcacc 240
cagggaaggg cctggagtgg attgggtaca tctattacag tgggagcacc tactacaacc 300
cgtccctcaa gagtcgagtt accatatcag tagacacgtc taagaaccag ttctccctga 360
agctgagctc tgtgactgcc gcgacacgg ccgtgtatta ctgtgcgaga gagcatctct 420
cctacggtga ctcgagatac tactactacg gtatggacgt ctggggccaa gggacccggt 480
caccgtctcc tcagcatccc egaccagccc caaggctctc ccgctgagcc tctgcagcac 540
ccagccagat gggaaacgtg tcatcgctg cctgggtccag ggcttcttcc ccaggagcc 600
actcagtgtg acctggagcg aaagggacag ggcgtgaccg ccagaaactt cccaccagc 660
caggatgcct ccggggacct gtacaccacg agcagccagc tgaccctgcc ggccacacag 720
tgcttagccg gcaagtccgt gacatgccac gtgaagcact acacgaatcc cagccaggat 780
gtgactgtgc cctgcccagt tccctcaact ccacctacce catctccctc aactccacct 840
accccatctc cctcatgctg ccacccccga ctgtcactgc accgaccggc cctcaggagc 900

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ctgctcttag gttcagaagc gaacctcagc tgcacactga ccggcctgag agatgctca 960
 ggtgtcacct tcacctggac gccctcaagt ggaagagcg ctgttcaagg accacctgag1020
 5 cgtgacctct gtggctgcta cagcgtgtcc agtgtcctgc cgggtgtgac cgagccatgg1080
 aaccatggga agaccttcac ttgcaactgct gctaccaccg agtccaagac cccgctaacc1140
 gccacctctt caaaatccgg aaacacattc cggcccgagg tccacctgct gccgccgccc1200
 tcggaggagc tggccctgaa cgagctggtg acgctgacgt gcctggcacg cggcttcagc1260
 cccaaggagc tgtctggttc ctggtctgag ggttcacagg agctgccccg cgagaagtac1320
 10 ctgacttggg catcccggca ggagcccagc cagggcacca ccaccttcgc tgtgaccagc1380
 atactgcgcg tggcagccga ggactggaag aagggggaca ctttctcctg catggtgggc1440
 cacgaggccc tggcgtggc cttcacacag aagaccatcg accgcttggc gggtaaacccl500
 acccatgtca atgtgtctgt tgtcatggcg gaggtggacg gcacctgcta ctgagccgcc1560
 cgctgtccc caccctgaa taaactccat gctcccccaa gcaaaaaaaa aaaaaaaa1620
 15 aaaaaaaaaa aaaaaaaaaa aaaa 1644

(2) INFORMATION ÜBER SEQ ID NO: 47:

20

(i) SEQUENZ CHARAKTERISTIK:

25

(A) LÄNGE: 1133 Basenpaare

(B) TYP: Nukleinsäure

30

(C) STRANG: einzel

(D) TOPOLOGIE: linear

35

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

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(iii) HYPOTHETISCH: NEIN

45

(iii) ANTI-SENSE: NEIN

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(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

55

(C) ORGAN:

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(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

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(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 47

atttatctgg gacagacatc ttcagaatga cacatgccaa acagtgggtc ttattaaatc 60
 aaagggttcag atattatcag attcagaaat agtgaatgctt tgtgtatcta ttttcttctc 120
 ttttaaacaga aaaagacaaa tgaatgggga aagacaatca ttgaatacaa aacaaataag 180
 ccatcacgcc tggccttctc tgatattgca cttttggaca tcgggtggtc tgaccaggaa 240
 ttcttttggtg acattggccc agtctgtttc aaataaatga actcaatcta aattaaaaaa 300
 55 gaaagaaatt tgaaaaaaot ttctctttgc catttcttct tcttcttttt taactgaaag 360
 ctgaatcctt ccatttcttc tgcacatcta cttgcttaaa ttgtgggcaa aagagaaaaa 420
 gaaggattga tcagagcatt gtgcaataca gtttcattaa ctcttcccc cgtccccca 480
 aaaatttgaa ttttttttcc aacactctta cactgtttat ggaaaatgtc aacctttgta 540
 agaaaaccaa aataaaaaat gaaaaataaa aaccataaac atttgacca cttgtggctt 600
 60 ttgaatatct tccacagagg gaagttaaa acccaaactt ccaaagggtt aaactacctc 660
 aaaacacttt cccatgagtg tgatccacat tgtaggtgac tgacctagac agagatgaac 720
 tgagggtcctt gttttgtttt gttcataata caaagggtgct aattaatagt atttcagata 780
 cttgaagaat gttgatggtg ctgagaagaat ttgagaagaa atactcctgt attgagttgt 840
 atcgtgtggt gtatttttta aaaaatttga ttttagcattc atattttcca tcttattccc 900
 65 aattaaaagt atgcagatta tttgcccata tcttcttcag attcagcatt tgttctttgc 960
 cagtctgatt ttcactctct tccatgggtc cacagaagct ttgtttcttg ggcaagcaga1020
 aaaattaaat tgtacctatt ttgtatatgt gagatgttta aataaattgt gaaaaaaatg1080
 aaataaagca tgtttggttt tccaaaagaa aaaaaaaaaa aaaaaagtcg acc 1133

(2) INFORMATION ÜBER SEQ ID NO: 48:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 969 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH
 (C) ORGAN:

(vii) SONSTIGE HERKUNFT:

- (A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 48

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gaggaggagg gtgtatctcc ttctgtcgga ccgccccttg gcttctgcac tgatggtggg 60
tggatgagta atgcatccag gaagcctgga ggctgtggt ttccgcaccc gctgccaccc120
ccgccctag cgtggacatt tatcctctag cgtcaggcc ctgccccat cgcgcagat180
ccagcgcca gagagacacc agagaacca ccatggcccc ctttgagccc ctggcttctg240
gcatcctgtt gttgctgtgg ctgatagccc ccagcagggc ctgcacctgt gtcccacccc300
accacagac ggccttctgc aattccgacc tcgtcatcag ggccaagttc gtggggacac360
cagaagtcaa ccagaccacc ttataccagc gttatgagat caagatgacc aagatgtata420
aagggttcca agccttaggg gatgccgtg acatccggtt cgtctacacc ccgcctatgg480
agagtgtctg cggatacttc cacagggtccc acaaccgcag cgaggagttt ctcatgtctg540
gaaaactgca ggatggactc ttgcacatca ctacctgcag ttctgtggct ccctggaaca600
gectgagctt agctcagcgc cggggcttca ccaagacctt cactgttggc tgtgaggaa660
gcacagtgtt tccctgttta tccatccctt gcaaaactgca gattggcaat cattgcttgt720
ggacggacca gtcctcccaa ggctctgaaa agggcttcca gtcccgtcac cttgcctgcc780
tgctcgggga gccagggtg tgcaactggc agtccctgcg gtcccagata gcctgaatcc840
tgcccggagt ggaagctgaa gcctgcaag tgccaccct gtcccactc ccatctttct900
tccggacaat gaaataaaga gttaccaccc agcaaaaaaa aaaaaaaaaa acaagtcgtc960
gcgtgctgt 969
  
```

(2) INFORMATION ÜBER SEQ ID NO: 49:

(i) SEQUENZ CHARAKTERISTIK:

- (A) LÄNGE: 617 Basenpaare
 (B) TYP: Nukleinsäure
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung
 hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 49

```
cctacaccta cctccctttt gggtttctat tcggaccgcg atgatttgct ttggaaggct 60
taacccccctt cttccccaaa cttgcccccg gagaaccccc agccttacga ccctcctcct120
gaagatgcaa aaccagcttg ccggccgcgc tctcttccag gacatcaaga agccagctga180
agatgagtgg ggtaaaaccc cagacgccat gaaagctgcc atggccctgg agaaaaagct240
gaaccagggc cttttggatc ttcatgccct gggttctgcc cgcacggacc cccatctctg300
tgacttcctg gagactcact tcctagatga ggaagtgaag ctatcaaga agatgggtga360
ccacctgacc aacctccaca ggctgggtgg ccgaggagct gggctgggag agtatctctt420
cgaaaggctc actctcaagc acgactaaga gccttctgag ccagcgact tctgaagggc480
cccttgcaaa gtaatagggc ttctgcctaa gcctctcctt ccagccaata ggcagctttc540
ttaactatcc taacaagcct tggaccaaat ggaaataaag ctttttgatg caaaaaaaga600
ggagggggga aaaaagc                                     617
```

(2) INFORMATION ÜBER SEQ ID NO: 50:

(i) SEQUENZ CHARAKTERISTIK:

(A) LÄNGE: 704 Basenpaare

(B) TYP: Nukleinsäure

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: aus einzelnen ESTs durch Assemblierung und Editierung hergestellte partielle cDNA

(iii) HYPOTHETISCH: NEIN

(iii) ANTI-SENSE: NEIN

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(C) ORGAN:

(vii) SONSTIGE HERKUNFT:

(A) BIBLIOTHEK: cDNA library

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO: 50

```
ggggagactc gtcaccaggc gtgcagtggg cactgctggg ctcccccatc ccgtcctaac 60
ccggaacagc cccgggcagg aggcgtggaa agtcgagggg gtaaaccgcg aatgtgcgtt120
gtgtaagcca cggcgaggc tggggcgagg gcgggacttg ggcgggcggg gtgggcttgg180
ccgagctggc ctccggggca ccgaccgcta taaggccagt cggactgcga cacagcccat240
cccctcgacc gctcgcgtcg catttgccg cctccctacc gctccaagcc cagccctcag300
ccatggcatg cccctggat caggccattg gcctcctcgt ggccatcttc cacaagtact360
```

cgggcagggg ggggtgacaag cacaccctga gcaagaagga gctgaaggag ctgatccaga420
 aggagctcac cattggctcg aagctgcagg atgctgaat tgcaaggctg atggaagact480
 tggaccggaa caaggaccag gaggtgaact tccaggagta tgtcaccttc ctgggggcct540
 tggctttgat ctacaatgaa gccctcaagg gctgaaaata aatagggag atggagacac600
 cctctggggg tctctctga gtcaaacca gtgggtgggt attgtacaat aaatttttt660
 tgggtcaaatt taaaaaaaa aaaaaaagg aaaaaagggt gagc 704

(2) INFORMATION ÜBER SEQ ID NO: 51:

- (A) LÄNGE: 95 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 51:

PCSSQFHPVE NRSQEPLAGD SMSPTLPVQ NMNNAMFLQK TLSLSFIGGN HQTAECTRL60
 SRTTDLSPSH SPYHHKSHNK KEKRYFGFKK SKKIM 95

(2) INFORMATION ÜBER SEQ ID NO: 52:

- (A) LÄNGE: 76 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 52:

LPRDTWFKLK CLTDHSRHLV HSYVNVSHLT WVHCLQTEHR LPLAWFENRN RAMPTDPSYV60
 WASKWNCTFI QIFTCL 76

(2) INFORMATION ÜBER SEQ ID NO: 53:

- (A) LÄNGE: 90 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 53:

RVNNCQEQLV VLKYNPQPRI PPVLQMDQLK QANTEDTKNE VRFIETRVTP LDELNTKMTL60
 TLSRYRSSET CLQNEIPEEF CSYPEIRGSN 90

(2) INFORMATION ÜBER SEQ ID NO: 54:

(A) LÄNGE: 117 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 54:

PLQPPRAMAP RGCIVAVFAI FCISRLCSH GAPVAPMTPY LMLCQPHKRC GDKFYDPLQH 60
 CCYDDAVVPL ARTQTCGNCT FRVCFEQCCP WTFMVKLINQ NCDSARTSDD RLCRSVS 117

(2) INFORMATION ÜBER SEQ ID NO: 55:

(A) LÄNGE: 103 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 55:

RVPQPALGWC RVDVGHRGHQ EGSESLGPHQ HTHMLSRIL EGDWASSGQ RQGGPQTGHR 60
 MKWAVECVFL WFPNSHSASQ ISGNTSLFLQ AHPGRIQES SFP 103

(2) INFORMATION ÜBER SEQ ID NO: 56:

- (A) LÄNGE: 81 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: ORF

10

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

15

(A) ORGANISMUS: MENSCH

:

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 56:

20

RCSFHTSGSW PRARRHHHSN SAAGGRRTCP HISCVAGTAS GKESWGPLGL RVSRGAWRCR60
KWQRQLRCSL GEPWLWVVAV E 81

(2) INFORMATION ÜBER SEQ ID NO: 57:

25

- (A) LÄNGE: 125 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

30

(ii) MOLEKÜLTYP: ORF

35

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

40

(A) ORGANISMUS: MENSCH

:

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 57:

45

RAARADSARA FPLPACKVVV PQGPPPGHVG AAGQAFPSFE RGFRCSRAS GLRSSLPSEF 60
SVVASPPPTH QSRCLGRAL GAMAPRGRKR KAEAAVVAVA EKREKLANGG EGMEEATVVI120
EHCTS 125

(2) INFORMATION ÜBER SEQ ID NO: 58:

50

- (A) LÄNGE: 119 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

55

(ii) MOLEKÜLTYP: ORF

60

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

65

(A) ORGANISMUS: MENSCH

:

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 58:

5 QRSPPFPFPP RSPASLASRL RRP RPQPYA SSRGEPWRLE PGRCSGTGG WGAETRPLSG 60
NWATKSAARK LCSYSGNLSQ RKGKLGPHQHP RGLADLGAQ PLCKQGAGRL EPNRLERLE 119

(2) INFORMATION ÜBER SEQ ID NO: 59:

- 10 (A) LÄNGE: 128 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
15 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

20 (iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

25 (A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 59:

30 TRQLVQCSIT TVASSIPSPP FASFSRFSAT ATTAASALRF LPRGAMAPRA RPRMQRDWWV 60
GGGDATERK LGNEERSPEA LLLQRKPLSK EGKAWPAAPT WPGGGPWGTT TLQAGSGKAR120
AESARAAR 128

(2) INFORMATION ÜBER SEQ ID NO: 60:

- 35 (A) LÄNGE: 127 Aminosäuren
(B) TYP: Protein
40 (C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

45 (iii) HYPOTHETISCH: ja

50 (vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 60:

55 VTVMQNFEL SFKYVLYSSY SWLKLDHTIA DCMVFTWTPC RMLDYLYSSY ANMLWAGEMK 60
SSSHQDLLFK WLDNWATKEL ELHLLGFELF WNTLLHFGKS KSSASGALSI ENLPSFALKD120
VLEFFIYT 127

(2) INFORMATION ÜBER SEQ ID NO: 61:

- 65 (A) LÄNGE: 111 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 61:

SIGPICSQGL GPGGIPSPIT LIKNGCNCKN PCLIIYLQLCS HLQMYLLMLS CQVPMQRWRG 60
LPLCGWGLWV VVKDRYQKNA FKCTNLLINI RCLLKKKKKK KKRVGCVGCI G 111

(2) INFORMATION ÜBER SEQ ID NO: 62:

(A) LÄNGE: 68 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 62:

YRSFTTTHKP HPHKGSPRHL CIGTWQLSIR RYICKWEHSC KYIRQGFLQL QPFLIKVIGE60
GIPPGPRP 68

(2) INFORMATION ÜBER SEQ ID NO: 63:

(A) LÄNGE: 195 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 63:

LVQPGGSCSG GRLLGVEFPS APRVRPFERS APAPATSLLG AMTTTTTFKG VDPNSRNSSR 60
VLRPPGGGSN FSLGFDEPTE QPVRKNKMAS NIFGTPEENQ ASWAKSAGAK SSGGREDLES120
SGLQRRNSSE ASSGDFDLK GEGDIHENVD TDLPGSLGQS EEKPVPAAPV PSPVAPAPVP180

SRRNPPGGKS SLVLG

(2) INFORMATION ÜBER SEQ ID NO: 64:

- (A) LÄNGE: 164 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 64:

VQSFPNSLL LENTHAMHR PKSQQRETC SSKEKKRQQ YIKCFFLMKQ IQEMYSQAQV 60
 VQFTSMEEED RTTAFRTVRA NRRGWTCRQ GDFFWMALGP GPPGWAQAQQ ARASLHSAPG120
 CLASLCPHFH EYHLLPSDLR SLRSLQRSS FSAVQMTPSL PCHH 164

(2) INFORMATION ÜBER SEQ ID NO: 65:

- (A) LÄNGE: 106 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 65:

FQAVSLYIQA FLCVRKALL ISQPVLLLSG YFLRLKNKRO FLCFAGGKAG GAGLFIVHMS 60
 QEEALSKGHW QVRATPRRLC GETPCGLGPG RRGACGLEMV CPVEAW 106

(2) INFORMATION ÜBER SEQ ID NO: 66:

- (A) LÄNGE: 349 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 66:

AALRS DAGMK RALGRRKGVW LRLRKILFCV LGLYIAIPFL IKLCPGIQAK LIFLNFVRVP 60
YFIDLKPKQD QGLNHTCNY LQPEEDVTIG VVHTVPAVWW KNAQKQDQMW YEDALASSHP120
IILYLHGNAG TRGGDHRVEL YKVLSSLGYH VVTFDYRGWG DSVGTPSERG MTYDALHVFD180
WIKARSGDNP VYIWGHS LGT GVATNLVRR L CERETPPDAL ILESPFTNIR EEAKSHPFVS240
IYRYFPGFDW FFLDPITSSG IKFANDENVK HISCPLILH AEDDPVVPFQ LGRKLYSIAA300
PARSFRDFKV QFVPFHS DLG YRHKYIYKSP ELPRILREFL GKSEPEHQH 349

(2) INFORMATION ÜBER SEQ ID NO: 67:

(A) LÄNGE: 191 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 67:

SGLSRLGPGR NQHAGQDVLC EVAAALHQVL KELLGQGIDY EKILKLTADA KFESGDVKAT 60
VAVLSFILSS AAKHSVDGES LSSELQQLGL PKEHAASLCR CYEEKQSPLQ KHLRVCSLRM120
NRLAGVGWRV DYTLS SLLQ SVEEPMVHLR LEVAAAPGTP AQPVAMSLA DKFQVLLAEL180
KQAQTLMSL G 191

(2) INFORMATION ÜBER SEQ ID NO: 68:

(A) LÄNGE: 164 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 68:

FFFFFFFFF FFFFSLLYFC LFFLLMKTAN NCLSREGKVM LGKVLRSPEP SSQERSEAAG 60
DLGGQSPGQG LSILEPGLPP EEQFRGRDSI RAGRLHTGLE HPSPQPRELI RVWACFSSAR120

RTWNLSAERD MATGWAGVPG AAATSSRRCT MGSSTDCSRL ELRV

164

(2) INFORMATION ÜBER SEQ ID NO: 69:

- (A) LÄNGE: 155 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 69:

NQGCLPKSSS EGVTPYGQGG STQAWNTLLL SPGSSSGSGP ASVLPGGPGT CLLRGTWQQA 60
 GLGSLGQLPP PAAGAPWALP RIAAGWSSGC SPPASPHLPT YSCVGCRPAS ASARGFASPH120
 NSGTGWPRAL WAAPAAVHW TRIRHRHCAW PHWRG 155

(2) INFORMATION ÜBER SEQ ID NO: 70:

- (A) LÄNGE: 35 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 70:

RRAAVTWVWL GVLCFESAVF TPTEVVRTCR LLRFS

35

(2) INFORMATION ÜBER SEQ ID NO: 71:

- (A) LÄNGE: 32 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

5

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 71:

KRLTQNTTPP TRPKSQLHVF KTSFKVSYFS TS

32

(2) INFORMATION ÜBER SEQ ID NO: 72:

10

(A) LÄNGE: 37 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

15

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

20

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

25

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 72:

ENRSNLHVL TSVGVKTADS KHNTFNQTQV TAARLQN

37

(2) INFORMATION ÜBER SEQ ID NO: 73:

35

(A) LÄNGE: 121 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

40

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

45

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

50

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 73:

LVKGMTVLEA VLEIQAITGS RLLSMVPGPA RPPGSCWDPT QCTRTWLLSH TPRRRWISGL 60
PRASCRLGEE PPPLPYCDQA YGEELSIRHR ETWAWLSRTD TAWPGAPGVK QARILGELLL120
V 121

55

(2) INFORMATION ÜBER SEQ ID NO: 74:

60

(A) LÄNGE: 115 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

65

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 74:

QACPWASLAQ GQTRLRRKL DTPVHGGLGL EGWLSGLEVP GGLPAGTRPS AAGWAVPCCC 60
CPQGLAVVAE DGTLSGWIRS PGSSSSRELR HKAGARLYTC RTQESLLQFL PEAPR 115

(2) INFORMATION ÜBER SEQ ID NO: 75:

(A) LÄNGE: 117 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 75:

RWHLIRLDQV TRQQQLSRAE AQGRGPAVHL QDPGEPVAVL ARSAEIASSV SLQQEQNQLW 60
PRWVGGS AFL AMAAATPRQE TAECLEGCNT RSNRQPPLFL MSDGQALQHL DRHGGWS 117

(2) INFORMATION ÜBER SEQ ID NO: 76:

(A) LÄNGE: 66 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 76:

PPQERRTIFV LYPRGSGREN MESGFYRLIG PIHKGHDWEK VWEQKENWDF RVQYAHPKLL 60
VAWGMS 66

(2) INFORMATION ÜBER SEQ ID NO: 77:

- (A) LÄNGE: 81 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: ORF

10

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

15

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 77:

20

ALSTRAMTGK RYGSRRRIGI LGCSTLTINF WWPGACLEAQ TVKQALLACL LVTTSAPAVL60
RLHPAPGTPP APEPPLSPCD G 81

(2) INFORMATION ÜBER SEQ ID NO: 78:

25

- (A) LÄNGE: 104 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

30

(ii) MOLEKÜLTYP: ORF

35

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

40

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 78:

45

TLLTIHIWTR DTVHEVHPSQ GDSGGSGAGG VPGAGWSLKT AGAEVVTSKQ ASRACLTVA 60
SRHAPGHQKF RVSVLHPKIP ILLLLPYLFP VMAVDRAYQ SIES 104

(2) INFORMATION ÜBER SEQ ID NO: 79:

50

- (A) LÄNGE: 104 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

55

(ii) MOLEKÜLTYP: ORF

60

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

65

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 79:

PSSPRAVRHS GHQDSKMASV VPVKDKLLE VKLGELPSWI LMRDFSPSGI FGAFQRGYYR 60
YNNKYINVKK GSISGITMVL ACYVLFYSYF SYKHLKHERL RKYH 104

(2) INFORMATION ÜBER SEQ ID NO: 80:

- (A) LÄNGE: 82 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 80:

RRGASRGLPW CWHATCSLAT PFPTSISSTS GSANTTEEDT LCTPPPHDLG PPSVVRNTIS60
IVAESFHILI GINLQIKHDW YV 82

(2) INFORMATION ÜBER SEQ ID NO: 81:

- (A) LÄNGE: 115 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

- (A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 81:

KDSATIEIVF LTEGLGPRSW GGGVQSVSS VVFAEPLVLE MLVGKGVAKE HVACQHHGNP 60
RDAPLLHIDV LVVVPVVTSL KRSENATRTE VPHQDPAWQL PQFDLQKFLV LHWYN 115

(2) INFORMATION ÜBER SEQ ID NO: 82:

- (A) LÄNGE: 187 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

- b) eine allelische Variation der unter a) genannten Nukleinsäure-Sequenzen
oder
c) eine Nukleinsäure-Sequenz, die komplementär zu den unter a) oder b) genannten Nukleinsäure-Sequenzen ist.
2. Eine Nukleinsäure-Sequenz gemäß einer der Sequenzen Seq. ID Nos. 1–50, oder eine komplementäre oder allelische Variante davon. 5
 3. Nukleinsäure-Sequenz Seq. ID No. 1 bis Seq. ID No. 50, dadurch gekennzeichnet, daß sie in Blasen-tumorge-webe erhöht exprimiert sind.
 4. BAC, PAC und Cosmid-Klone, enthaltend funktionelle Gene und ihre chromosomale Lokalisation, entsprechend den Sequenzen Seq. ID. No. 1 bis Seq. ID No. 50, zur Verwendung als Vehikel zum Gentransfer. 10
 5. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 4, dadurch gekennzeichnet, daß sie eine 90%ige Ho-mologie zu einer humanen Nukleinsäure-Sequenz aufweist.
 6. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 4, dadurch gekennzeichnet, daß sie eine 95%ige Ho-mologie zu einer humanen Nukleinsäure-Sequenz aufweist.
 7. Eine Nukleinsäure-Sequenz, umfassend einen Teil der in den Ansprüchen 1 bis 6 genannten Nukleinsäure-Se-quenzen, in solch einer ausreichenden Größe, daß sie mit den Sequenzen gemäß den Ansprüchen 1 bis 6 hybridisie-ren. 15
 8. Ein Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 7, dadurch gekennzeichnet, daß die Größe des Frag-ments eine Länge von mindestens 50 bis 4500 bp aufweist.
 9. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 7, dadurch gekennzeichnet, daß die Größe des Frag-ments eine Länge von mindestens 50 bis 4000 bp aufweist. 20
 10. Eine Nukleinsäure-Sequenz gemäß einem der Ansprüche 1 bis 9, die mindestens eine Teilsequenz eines biolo-gisch aktiven Polypeptids kodiert.
 11. Eine Expressionskassette, umfassend ein Nukleinsäure-Fragment oder eine Sequenz gemäß einem der Ansprü-che 1 bis 9, zusammen mit mindestens einer Kontroll- oder regulatorischen Sequenz. 25
 12. Eine Expressionskassette, umfassend ein Nukleinsäure-Fragment oder eine Sequenz gemäß Anspruch 11, wo-rin die Kontroll- oder regulatorische Sequenz ein geeigneter Promotor ist.
 13. Eine Expressionskassette gemäß einem der Ansprüche 11 und 12, dadurch gekennzeichnet, daß die auf der Kas-sette befindlichen DNA-Sequenzen ein Fusionsprotein kodieren, das ein bekanntes Protein und ein biologisch aktives Polypeptid-Fragment umfaßt. 30
 14. Verwendung der Nukleinsäure-Sequenzen gemäß den Ansprüchen 1 bis 10 zur Herstellung von Vollängen-Ge-nen.
 15. Ein DNA-Fragment, umfassend ein Gen, das aus der Verwendung gemäß Anspruch 14 erhältlich ist.
 16. Wirtszelle, enthaltend als heterologen Teil ihrer exprimierbaren genetischen Information ein Nukleinsäure-Fragment gemäß einem der Ansprüche 1 bis 10. 35
 17. Wirtszelle gemäß Anspruch 16, dadurch gekennzeichnet, daß es ein prokaryontisches oder eukaryontische Zell-system ist.
 18. Wirtszelle gemäß einem der Ansprüche 16 oder 17, dadurch gekennzeichnet, daß das prokaryontische Zellsy-tem E. coli und das eukaryontische Zellsystem ein tierisches, humanes oder Hefe-Zellsystem ist.
 19. Ein Verfahren zur Herstellung eines Polypeptids oder eines Fragments, dadurch gekennzeichnet, daß die Wirts-zellen gemäß den Ansprüchen 16 bis 18 kultiviert werden. 40
 20. Ein Antikörper, der gegen ein Polypeptid oder ein Fragment gerichtet ist, welches von den Nukleinsäuren der Sequenzen Seq. ID No. 1 bis Seq. ID No. 50 kodiert wird, das gemäß Anspruch 19 erhältlich ist.
 21. Ein Antikörper gemäß Anspruch 20, dadurch gekennzeichnet, daß er monoklonal ist.
 22. Ein Antikörper gemäß Anspruch 20 dadurch gekennzeichnet, daß er ein Phage-Display-Antikörper ist. 45
 23. Polypeptid-Teilsequenzen, gemäß den Sequenzen Seq. ID Nos. ORF 51–106.
 24. Polypeptid-Teilsequenzen gemäß Anspruch 22, mit mindestens 80%iger Homologie zu diesen Sequenzen.
 25. Ein aus einem Phage-Display hervorgegangenen Polypeptid, welches an die Polypeptid-Teilsequenzen gemäß Anspruch 24 binden kann.
 26. Polypeptid-Teilsequenzen gemäß Anspruch 22, mit mindestens 90%iger Homologie zu diesen Sequenzen. 50
 27. Verwendung der Polypeptid-Teilsequenzen gemäß den Sequenzen Seq. ID No. 51–106, als Tools zum Auffin-den von Wirkstoffen gegen den Blasen-tumor.
 28. Verwendung der Nukleinsäure-Sequenzen gemäß den Sequenzen Seq. ID No. 1 bis Seq. ID No. 50 zur Expres-sion von Polypeptiden, die als Tools zum Auffinden von Wirkstoffen gegen den Blasen-tumor verwendet werden können. 55
 29. Verwendung der Nukleinsäure-Sequenzen Seq. ID No. 1 bis Seq. ID No. 50 in sense oder antisense Form.
 30. Verwendung der Polypeptid-Teilsequenzen Seq. ID No. 51–106 als Arzneimittel in der Gentherapie zur Be-handlung des Blasen-tumors.
 31. Verwendung der Polypeptid-Teilsequenzen Seq. ID No. 51–106, zur Herstellung eines Arzneimittels zur Be-handlung gegen den Blasen-tumor. 60
 32. Arzneimittel, enthaltend mindestens eine Polypeptid-Teilsequenz Seq. ID No. 51–106.
 33. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 10, dadurch gekennzeichnet, daß es eine genomische Sequenz ist.
 34. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 10, dadurch gekennzeichnet, daß es eine mRNA-Se-quenz ist. 65
 35. Genomische Gene, ihre Promotoren, Enhancer, Silencer, Exonstruktur, Intronstruktur und deren Spleißvarian-ten, erhältlich aus den cDNAs der Sequenzen Seq. ID No. 1 bis Seq. ID No. 50.
 36. Verwendung der genomischen Gene gemäß Anspruch 33, zusammen mit geeigneten regulativen Elementen.

37. Verwendung gemäß Anspruch 34, dadurch gekennzeichnet, daß das regulative Element ein geeigneter Promotor und/oder Enhancer ist.

38. Eine Nukleinsäure-Sequenz gemäß den Ansprüchen 1 bis 7, dadurch gekennzeichnet, daß die Größe des Fragments eine Länge von mindestens 300 bis 3500 bp aufweist.

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Hierzu 10 Seite(n) Zeichnungen

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Systematische Gen-Suche in der Incyte LifeSeq Datenbank

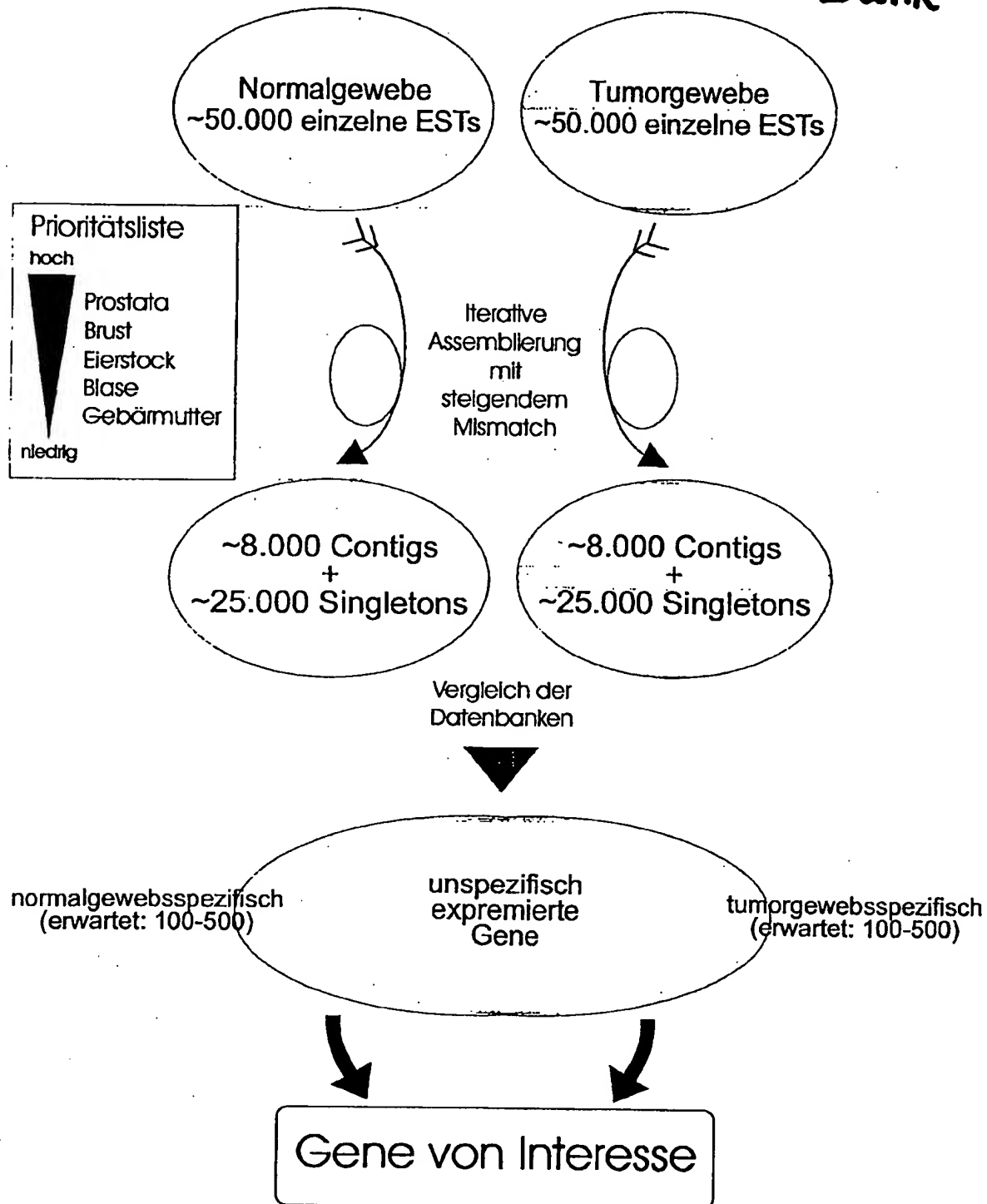


Fig. 1

Prinzip der EST-Assemblierung

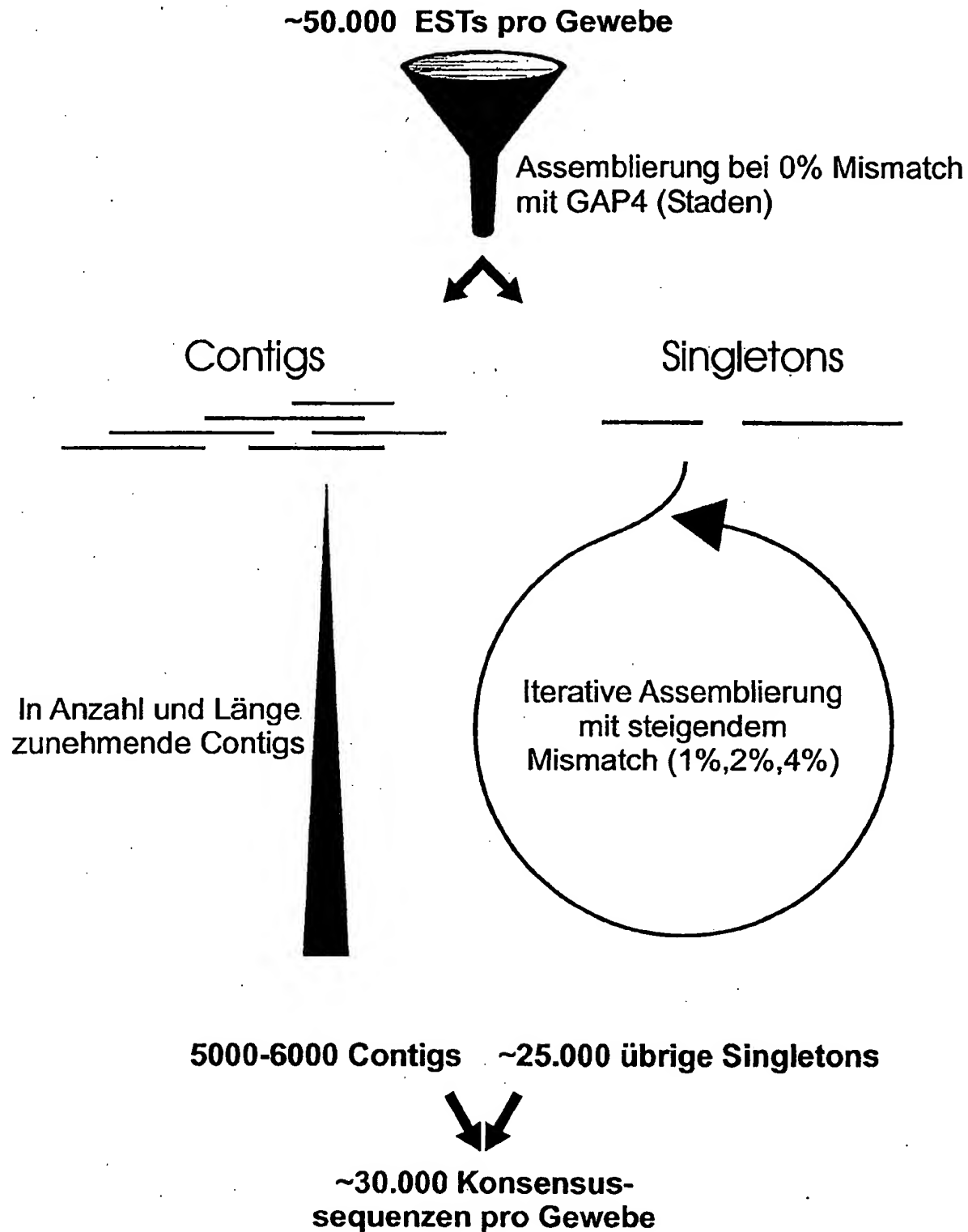


Fig. 2a

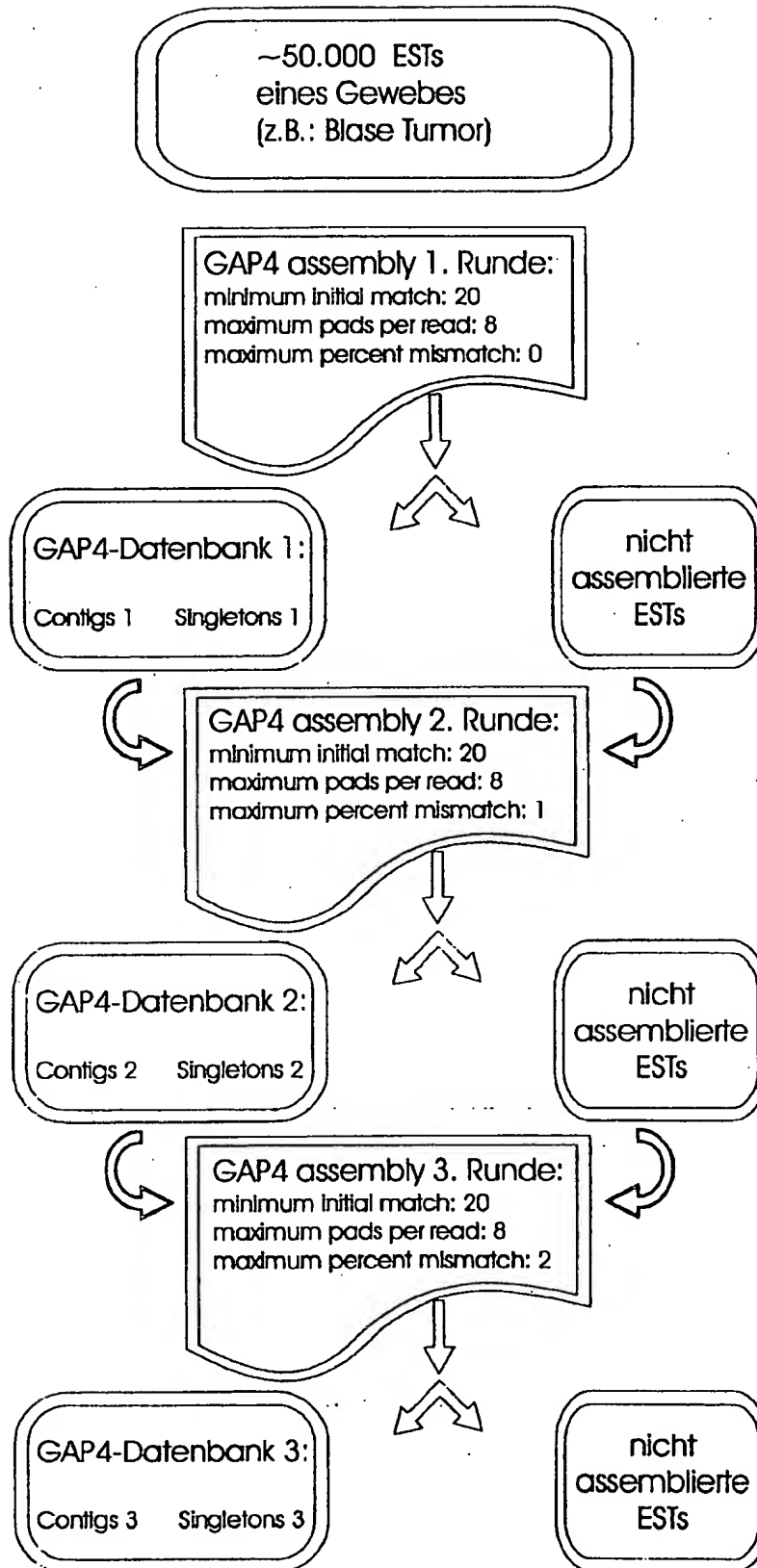


Fig. 2b1

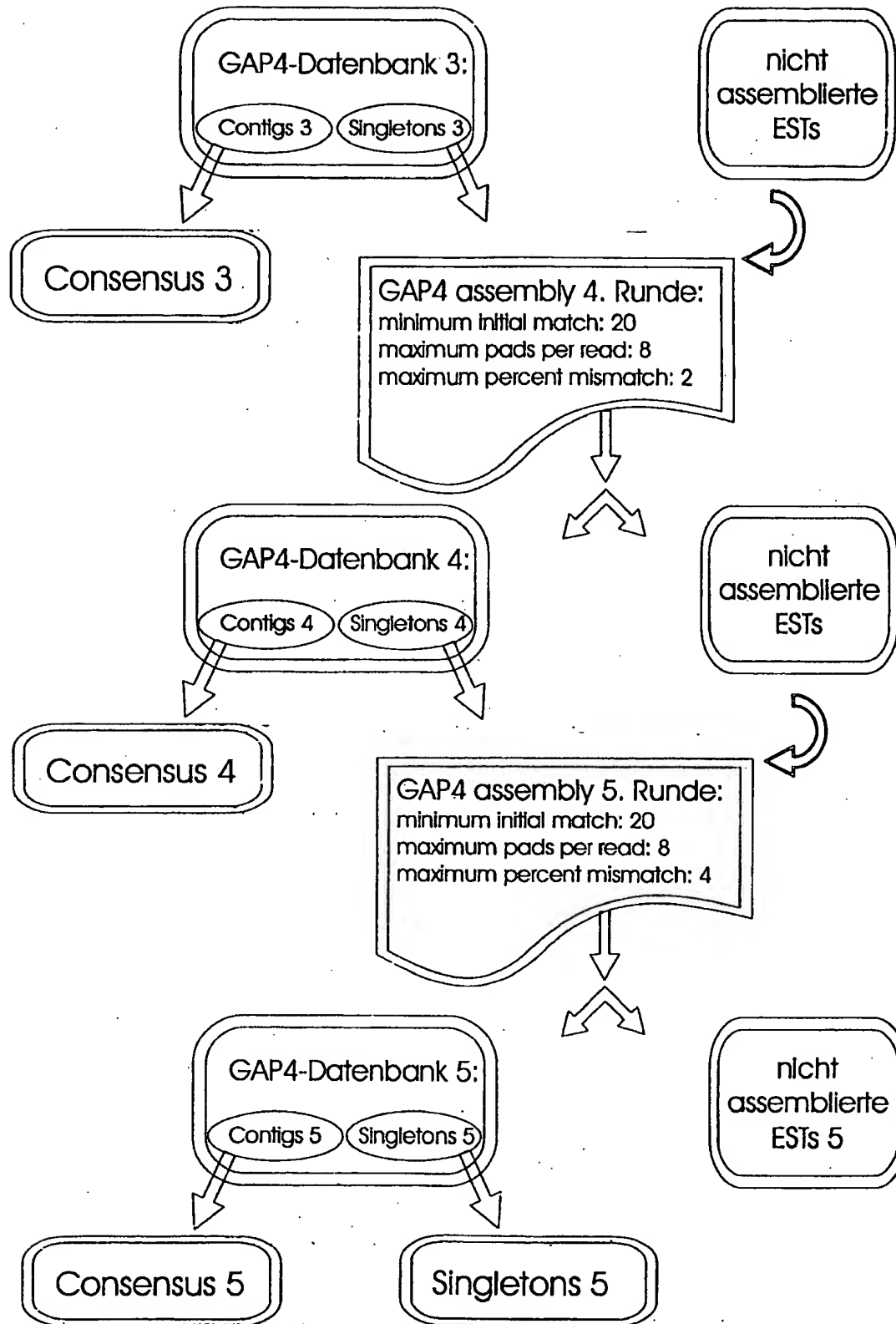


Fig. 2b2

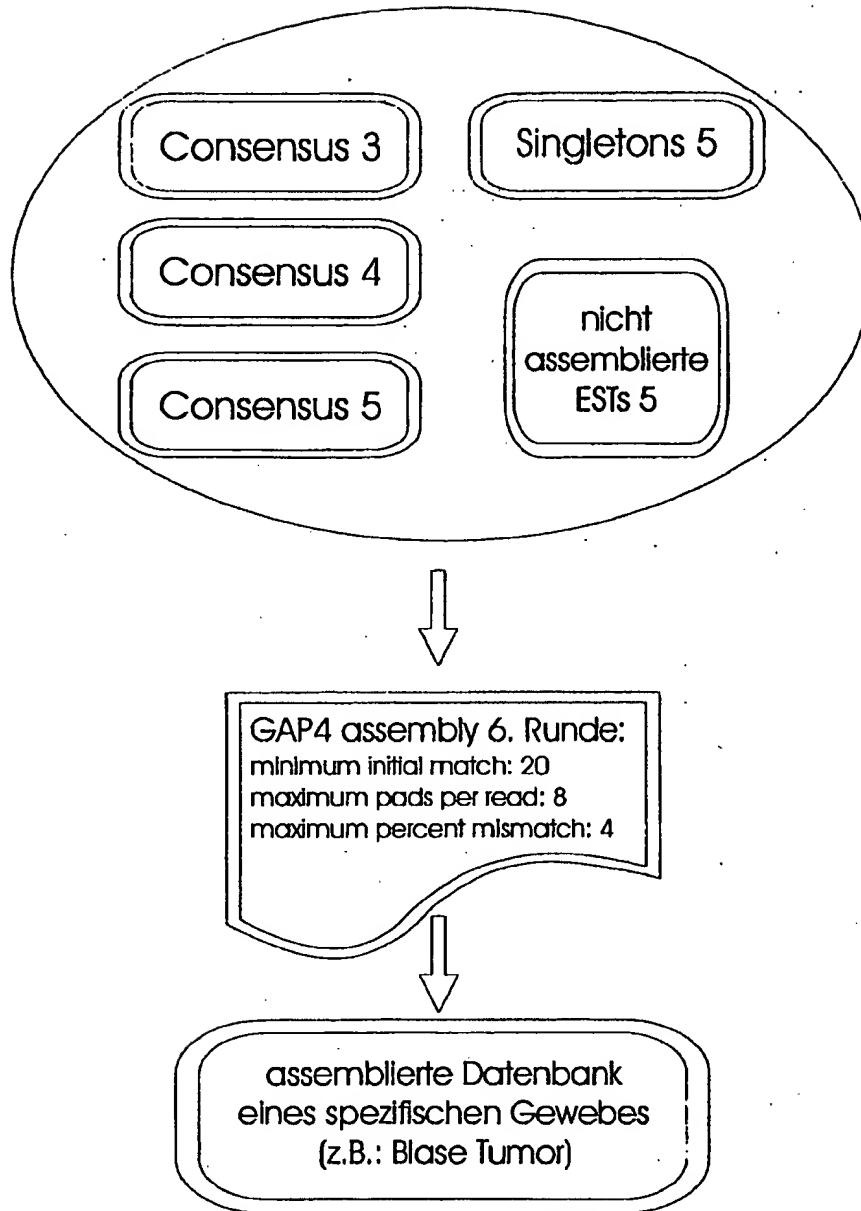


Fig. 2b3

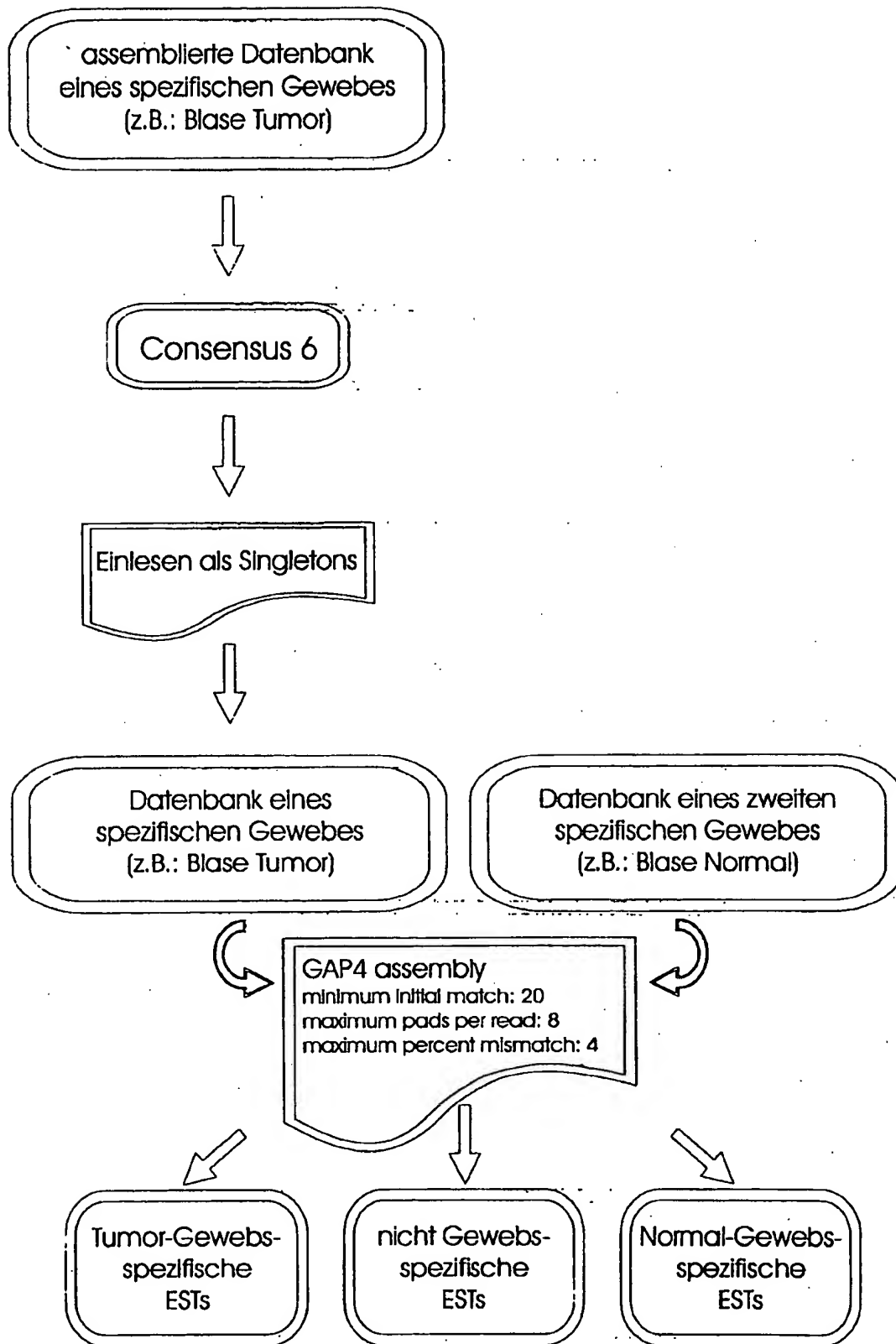


Fig. 2b4

In silico Subtraktion der Genexpression in verschiedenen Geweben

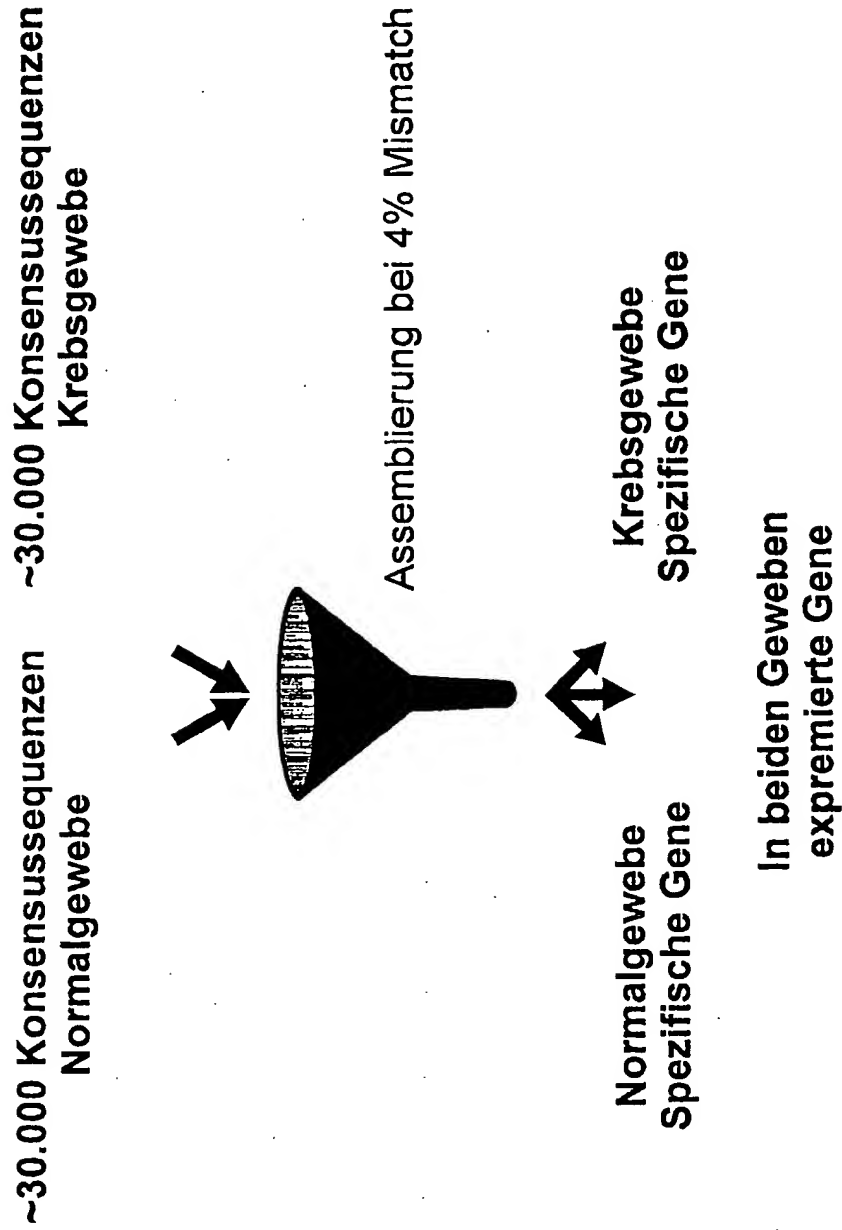


Fig. 3



Gene von Interesse



Bestimmung der
gewebsspezifischen
Expression über
elektronischen Northern
(INCYTE LifeSeq und
öffentliche EST
Datenbanken)



Kandidatengene für
Tumorsuppressoren oder
Tumoraktivatoren

Fig. 4a

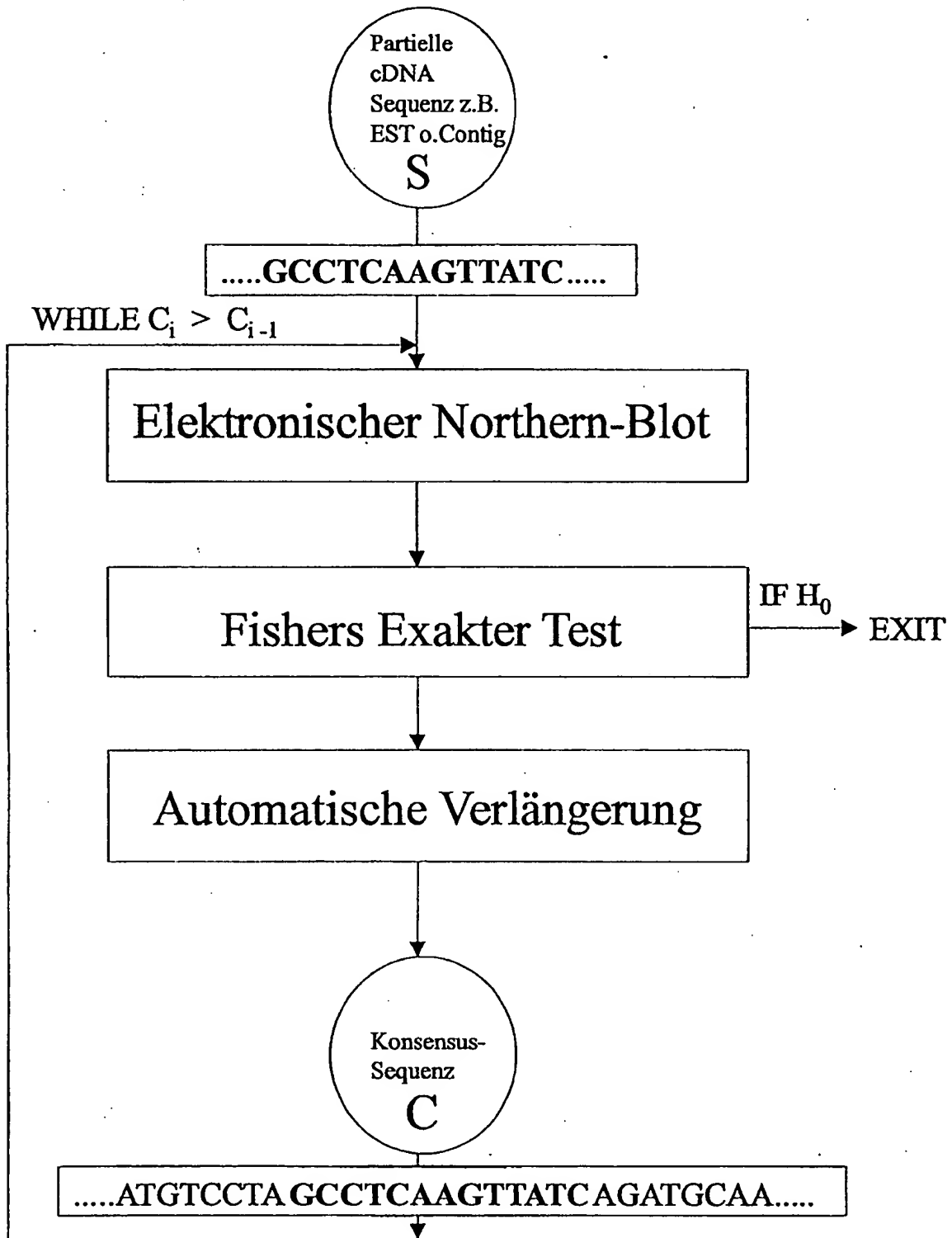


Fig. 4b

Isolieren von genomischen BAC und PAC Klonen

Chromosomale Klon-Lokalisation über FISH



Hybridisierungssignal



Sequenzierung von Klonen, die in Regionen lokalisiert sind, die chromosomale Deletionen in Prostata- und Brustkrebs aufweisen, führt zur Identifizierung von Kandidatengenen



Bestätigung der Kandidatengene durch Screening von Mutationen und/oder Deletionen in Krebsgeweben

Fig. 5

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 82:

ARAARGARRT SRAVTPTCAT PAGPMPCSLR PPSLRCSLHS ACCSGDPASY RLWGAPLQPT 60
 LGVVPQASVP LLTDLAQWEP VLVPEAHPNA SLTMYVCTPV PHPDPPMALS RTPTRQISSS120
 DTDPPADGPS NPLCCCFHGP AFSTLNPVLR HLFPPQEAFFA HPIYDLSQVW SVVSPAPSRG180
 QALRRAQ 187

(2) INFORMATION ÜBER SEQ ID NO: 83:

(A) LÄNGE: 241 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 83:

FFFFGLSNRC LLRAYAVLRL PFREPHECEA WPLPPGLQAP SLETPRNSRR LLSSSSTQST 60
 SSQPLLGPPE CLSPAGCGGH HGPDLAQVID GVGREGFLGE EVPEHRVKGG ECWAMETAAE120
 RVGGAICRRI CVTRADLPGG SPGEGHGRVR VGHGADIHG QTCVRMCLRN QDRLPLGQVC180
 EEWHRGLGHH TQCGLRGPP EPIAGRVPR AGRVQGAAG WRQPAAGHGP RWRCTSRCHS240
 T 241

(2) INFORMATION ÜBER SEQ ID NO: 84:

(A) LÄNGE: 113 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 84:

MGWAGKASWG KRCLSTGLRV ENAGPWKQQQ RGLEGPSACG SVSLELICRV GVLERAMGGS 60
GWGTGVQTYM VRLALGCASG TRTGSHWARS VRSGTEAWGT TPSVGOSGAP QSL 113

(2) INFORMATION ÜBER SEQ ID NO: 85:

- (A) LÄNGE: 107 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 85:

AFLSFLFSER FKASTTLEFP SLLNLICTKS FALVGVVETA LSLSTSVREC EPPWQVPVQG 60
PAALHLGRVT GAPAVCPKAS PWPFGLSLGR FRTEHQGRQA FQGISIN 107

(2) INFORMATION ÜBER SEQ ID NO: 86:

- (A) LÄNGE: 107 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 86:

LRNRLWKVKL EEPDLISPTS KTPSEQRPQ HPPRTGDSIF MATPCGGRLT TSHHIPELS 60
SSSGMTPSPP PPSSSFSSFC LEVSELSCLS FFLRDSKPPR LCFPRPF 107

(2) INFORMATION ÜBER SEQ ID NO: 87:

- (A) LÄNGE: 115 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

5

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 87:

IQKGRGKQSR GGFESLRKKE RQESSETKRQ KDEKEEEGGG GDGVIPLEEL SSGMMWWLVV 60
NRPPQGVAMK MESPVRRGCC GRCCSDGVFD VGLMRSGSSS FTFQSRFLSQ VVQWL 115

10

(2) INFORMATION ÜBER SEQ ID NO: 88:

(A) LÄNGE: 124 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

15

(ii) MOLEKÜLTYP: ORF

20

(iii) HYPOTHETISCH: ja

25

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

30

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 88:

CSVRNLPRLR PKQGGEAFGH TAGAPVTRPR WRAAGPCTGT CQGGSHSRTL VLRLKAVSTT 60
PTSANDLVQI KFRRDGGNRV VEALNLSEKR KDRKAQKQRD RRMKKKKREV VGTASSRWRS120
SALG 124

35

(2) INFORMATION ÜBER SEQ ID NO: 89:

(A) LÄNGE: 198 Aminosäuren
(B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

40

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

50

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

55

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 89:

EGAGGEWRCP AAGGPRGEDG PPGLRLTERA GLTTRHLTGT ADPSQKHNL R DPSSSKDERI 60
QQTQPNREGA CSLNLHGLRY LCKPGVAPVL RVILRSCLFP NGFTSGSCRF SLGLSLILKW120
VAGGRGWL RP LLALRLQSWE QDTSPEFHFF SCPNHAHTIV QNQSTFEKWL HGHPPGPRKL180
HSKGLFTWQQ NPSPAVSP 198

60

65

(2) INFORMATION ÜBER SEQ ID NO: 90:

- (A) LÄNGE: 124 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 90:

LPPVEPPVLK GSCRKDMHRN EERERSENEV WRARPGPTAQ GSSPPPDAPF HPPPQCLLSP 60
KAPASEVLGA HPPQCGQGK GQVLDLAKCP EMTLLLTTHFF GPWQSPTCPQ HGAPGRTGRQ120
EGGW 124

(2) INFORMATION ÜBER SEQ ID NO: 91:

- (A) LÄNGE: 147 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 91:

NSGEVSCSQL CSLRASGRS HPLPPATHFK MRLRPRLKRQ LPEVNPFGKR HERRMTLRTG 60
ATPGLHKYRR PWRLRLQAPS LLGCVCCILS SLELLGSLRL CFWDGSAVPV RCLVVRPALS120
VSLSPGGPSS PLGPPAAGHL HSPPAPS 147

(2) INFORMATION ÜBER SEQ ID NO: 92:

- (A) LÄNGE: 374 Aminosäuren
- (B) TYP: Protein
- (C) STRANG: einzel
- (D) TOPOLOGIE: linear

(II) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 92:

SREAPESRRW AVWRSLES LP RHQLCLPVG APPAPAMLSA LARPASAALR RSFSTSAQNN 60
 AKVAVLGASG GIGQPLSLLL KNSPLVSRLT LYDIAHTPGV AADLSHIETK AAVKGYLGPE120
 QLPDCLKGCD VVIPAGVPR KPGMTRDDL F NTNATIVATL TAACAQHCPE AMICVIANPV180
 NSTIPITAEV FKKHGVYNPN KIFGVTTLDI VRANTFVAEL KGLDPARVNV PVIGGHAGKT240
 IIP LISQCTP KVD FPQDQLT ALTGRIQEAG TEVV KAKAGA GSATLSMAYA GARFVFS LVD300
 AMNGKEGVVE CSFVKSQETE CTYFSTPLLL GKKGIEKNLG IGKVSSFEEK MISDAIPELK360
 ASI KKGEDFV KTLK 374

(2) INFORMATION ÜBER SEQ ID NO: 93:

(A) LÄNGE: 238 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 93:

LNEGTFHNTF LSIHCHKGE DKAGAGIRHG EGGRACSGFS LDHLRAGLLD PPSECCQLVL 60
 GKVHLGGALR DQGDDGLPSM ATNDRD VDSS WIQTLQLCNK GVGSDDVQGR HAEDFVG VVH120
 SMLLENFCCD GDGGINRIGN DADHGFR AVL GTGSGQGGHN RGIGVEQVVP GHAWLSGDSS180
 RNNYHITTFQ AVRQLERSEV AFHSGFGLDV AQICGHSGCV RDIIEGQAAH QGAVLQEK 238

(2) INFORMATION ÜBER SEQ ID NO: 94:

(A) LÄNGE: 242 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 94:

EAGEEKAEEE GVAEEGVNK FSYPSSHREC CPAVEEEDDE EAVKKEAHR TSTSALSPGSK 60
 PSTWVSCPG EENQATEDKR TERSKGARKT SVSPRSSGSD PRSWEYRSGE ASEEEK EKAH120
 KETGKGEAAP GPQSSAPAQR PQLKSWWCQP SDEEEGEVKA LGAAEKDGEA ECPPCI PPPS180
 AFLKAWVYWP GEDTEEEDE EDEDS DSGS DEEEGEAEAS SSTPATGVFL KSWVYQPGED240
 TQ 242

(2) INFORMATION ÜBER SEQ ID NO: 95:

(A) LÄNGE: 237 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 95:

RPGCIGQERT LDSEDKEDDS EAALGEAESD PHPSHPDQRA HFRGWGYRPG KETEEEEAAE 60
 DWGEAEPCPF RVAIYVPEK PPPPWAPPRL PLRLQRRLLR PETPTHDPDP ETPLKARKVR120
 FSEKVTVHFL AVWAGPAQAA RQGPWEQLAR DRSRFARRIT QAQEELSPCL TPAARARAWA180
 RLRNPPLAPI PALTQTLPS SVPSSPVQTT PLSQAVATPS RSSAAAAAAL DLSGRRG 237

(2) INFORMATION ÜBER SEQ ID NO: 96:

(A) LÄNGE: 890 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 96:

QDEHLITFFV PVFEPLPPQY FIRVVS DRWL SCETQLPVSE RHLILPEKYP PPTELLDLQP 60
 LPVSALRNSA FESLYQDKFP FENPIQTQVF NTVYNSDDNV FVGAPTGS GK TICAFAILR120
 MLLQSSEGR C VYITPMEALA EQVYMDWY EK FQDRLNKKV LITGETSTDL KLLGKGNII180
 STPEKWDILS RRWKQRKNVQ NINLFVVDEV HLIGGENGPV LEVICSRMRY ISSQIERPIR240
 IVALSSSLSN AKDVAHWLGC SATSTFNHFP NVRPVPLELH IQGFNISHTQ TRLLSMAKPV300
 YHAITKHSPK KPVIVFVPSR KQTRLTAIDI LTTCAADIQR QRFLHCTEKD LIPYLEKLS360
 STLKETLLNG VGYLHEGLSP MERRLVEQLF SSGAIQVVVA SRSLCWGMNV AAHLVIIMDT420
 QYNGKIHA Y VDYPIYDVLQ MVGHANRPLQ DDEGR CVIMC QGSKKDFFK FLYEPLPVES480
 HLDHCMHDH F NAEIVTKTIE NKQDAVDYLT WFLYRRMTQ NPNYYNLQGI SHRHLSDHLS540
 ELVEQTLSD L EQSKCISIED EMDVAPLN LG MIAAYYYINY TTIELFSMSL NAKTKVRGLI600
 EIISNAEYE NIPIRHHEDN LLRQLAQKVP HKLNNPKFND PHVKTNLLQ AHLSRMQLSA660
 ELQSDTEEIL SKAIRLIQAC VDLSSNGWL SPALAAMELA QMVTQAMWSK DSYLKQLPHF720
 TSEIRKRC TD KGVESVFDIM EMEDEERNAL LQLTDSQIAD VARFCNRYPN IELSYEVVDK780
 DSIRSGGPV VLVQLEREE VTGPVIAPLF PQKREEGWV VIGDAKSNL ISIKRLTLQ840
 KAKVKLDFVA PATGAHNYTL YFMSDAYMGC DQEKFSVDV KEAETDSDSD 890

(2) INFORMATION ÜBER SEQ ID NO: 97:

- (A) LÄNGE: 281 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: ORF

10

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

15

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 97:

20

GDGSAEHGPR PLAAPLVTSR GAPASARPRG ALPGGSAPSA PHGQLPGRAQ PAPVSGPPPT 60
 SGLCHFDPAW PWPLWPGPWQ LPPHPQDWPA QPDIPQDWVS FLRSFGQLTL CPRNGTVTGK120
 WRGSHVVGLL TTLNFGDGPD RNKTRTFQAT VLGSQMGLKG SSAGQLVLIT ARVTERTAG180
 TCLYFSAVPG ILPSSQPPIS CSEEGAGNAT LSPRMGEECV SVWSHEGLVL TKLLTSEELA240
 LCGSRLLVLG SFLLLFCGLL CCVTAMCFHP RRESHWSRTR L 281

25

(2) INFORMATION ÜBER SEQ ID NO: 98:

30

- (A) LÄNGE: 206 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

35

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

40

(vi) HERKUNFT:

45

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 98:

50

RLEPRSVTRS RRAVSRLSAR PGKVSAMAF LASGPYLTHQ QKVLRLYKRA LRHLESWCVQ 60
 RDKYRYFACL MRARFEEHKN EKDMAKATQL LKEAEEEFWY RQHPQPYIFP DSPGGTSYER120
 YDCYKVPWC LDDWHPSEKA MYPDYFAKRE QWKLLRRESW EREVQLQEE TPPGGPLTEA180
 LPPARKEGDL PPLWWYIVTR PRERPM 206

55

(2) INFORMATION ÜBER SEQ ID NO: 99:

- (A) LÄNGE: 139 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

60

(ii) MOLEKÜLTYP: ORF

65

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 99:

PLVPSFPSAV SSTVLSWQSN QDTLPSQKDA SHLSTILGPC SNRISHRRCP QESQGRCAV 60
DADGTRILPR PPSAAGWPSP YPFHSYVLQT GLSSNKQSIG ICLSGRTTTR GGVAPAYKAA120
TPFADGSGRV PTPRTPLRR 139

(2) INFORMATION ÜBER SEQ ID NO: 100:

(A) LÄNGE: 79 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 100:

APFWDLVVAIV SLIGGAPRRV REDVWLWMLT VPEFFLGLLQ QLGGLRHILF ILMFFKPGSH60
QTSKVSVEVS LDAPRLEVA 79

(2) INFORMATION ÜBER SEQ ID NO: 101:

(A) LÄNGE: 89 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:
(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 101:

VGGACAVALP QAAAMAGQED PVQREIHQDW ANREYIEIIT SSIKKIADFL NSFDMSCRSR60
LATLNEKLTA LERRIEYIEA RVTKGETLT 89

(2) INFORMATION ÜBER SEQ ID NO: 102:

(A) LÄNGE: 88 Aminosäuren

- (B) TYP: Protein
(C) STRANG: einzel
(D) TOPOLOGIE: linear

5

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

10

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

15

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 102:

NSAVLLLGSC FTQHRPHGKG PSSQLLPFS LKSNRAYSCF SFFKSVAFGL CHLGCGVVC60
KKFRGTVGND VRHETFSVTF YTSTCQCI 88

20

(2) INFORMATION ÜBER SEQ ID NO: 103:

25

(A) LÄNGE: 89 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

30

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

35

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

40

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 103:

HFIHLLVNVE ETFTAQSLGL FVKVLLTSIF LSLSLKLSLK FSLPLHCFCE QSLSPPHLWW60
EVRGSPGQDT HALAMWLPEN VVANPPVSC 89

45

(2) INFORMATION ÜBER SEQ ID NO: 104:

50

(A) LÄNGE: 240 Aminosäuren

(B) TYP: Protein

(C) STRANG: einzel

(D) TOPOLOGIE: linear

55

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

60

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

65

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 104:

5 REQILFIEIR DTAKGGETEQ PPSLSPLHGG RMPGEGEQ SLARETQSHR GRRQGWATW 60
 VTRCRESLNR GGAGAGKRAG ALAHHVFLAL IEPNLAEREA SEEEVKACSD ETVVADLLVK120
 VVYVLGAILK IFLREGNVLN QHSGMDIEKY SEHYQHDHSP GAEDDAAGGQ LRPTAQERRH180
 KEGSRGSPRC KRARKAVGES PGCPRGAEG AWPRQPSPH GAAGGKGWAP LRTFAGDGAV240

(2) INFORMATION ÜBER SEQ ID NO: 105:

- 10 (A) LÄNGE: 136 Aminosäuren
 (B) TYP: Protein
 15 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 105:

30 RLYMFWGPFS KSFSVKGTS INTAEWTLKN IPSIINMITP QALRTMPQAA SFGPQHRSDA 60
 IKKGVGDRRG ASGLGKRWEK AQDALAGGQR GRGPGLNHPI RGRQAEKAGL LSGLSRETAP120
 SETKTAPGET FLDLCS 136

(2) INFORMATION ÜBER SEQ ID NO: 106:

- 40 (A) LÄNGE: 173 Aminosäuren
 (B) TYP: Protein
 (C) STRANG: einzel
 (D) TOPOLOGIE: linear

(ii) MOLEKÜLTYP: ORF

(iii) HYPOTHETISCH: ja

(vi) HERKUNFT:

(A) ORGANISMUS: MENSCH

(xi) SEQUENZ-BESCHREIBUNG: SEQ ID NO 106:

60 LQSRKVSPG AVLVSDBGVS RESPERSPAF SACRPRMGWL RGPRLCPP ARASWAFSHR 60
 FPSPLAPRRS PTPFFMASLL CCGPKLAACG IVLSAWGVIM LIMLGIFNV HSAVLIEDVP120
 FTEKDFENGQ QNIYNLYEQV SYNCFIAAGL YLLGGFSFC QVRLNKRKEY MVR 173

Patentansprüche

- 65 1. Eine Nukleinsäure-Sequenz, die ein Genprodukt oder ein Teil davon kodiert, umfassend
 a) eine Nukleinsäure-Sequenz, ausgewählt aus der Gruppe Seq. ID No. 2-5, 7-13, 16, 18, 20, 23, 26-27,
 31-32, 36, 45.

Impact of DNA Amplification on Gene Expression Patterns in Breast Cancer^{1,2}

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ABSTRACT

Genetic changes underlie tumor progression and may lead to cancer-specific expression of critical genes. Over 1100 publications have described the use of comparative genomic hybridization (CGH) to analyze the pattern of copy number alterations in cancer, but very few of the genes affected are known. Here, we performed high-resolution CGH analysis on cDNA microarrays in breast cancer and directly compared copy number and mRNA expression levels of 13,824 genes to quantitate the impact of genomic changes on gene expression. We identified and mapped the boundaries of 24 independent amplicons, ranging in size from 0.2 to 12 Mb. Throughout the genome, both high- and low-level copy number changes had a substantial impact on gene expression, with 44% of the highly amplified genes showing overexpression and 10.5% of the highly overexpressed genes being amplified. Statistical analysis with random permutation tests identified 270 genes whose expression levels across 14 samples were systematically attributable to gene amplification. These included most previously described amplified genes in breast cancer and many novel targets for genomic alterations, including the *HOXB7* gene, the presence of which in a novel amplicon at 17q21.3 was validated in 10.2% of primary breast cancers and associated with poor patient prognosis. In conclusion, CGH on cDNA microarrays revealed hundreds of novel genes whose overexpression is attributable to gene amplification. These genes may provide insights to the clonal evolution and progression of breast cancer and highlight promising therapeutic targets.

INTRODUCTION

Gene expression patterns revealed by cDNA microarrays have facilitated classification of cancers into biologically distinct categories, some of which may explain the clinical behavior of the tumors (1-6). Despite this progress in diagnostic classification, the molecular mechanisms underlying gene expression patterns in cancer have remained elusive, and the utility of gene expression profiling in the identification of specific therapeutic targets remains limited.

Accumulation of genetic defects is thought to underlie the clonal evolution of cancer. Identification of the genes that mediate the effects of genetic changes may be important by highlighting transcripts that are actively involved in tumor progression. Such transcripts and their encoded proteins would be ideal targets for anticancer therapies, as demonstrated by the clinical success of new therapies against amplified oncogenes, such as *ERBB2* and *EGFR* (7, 8), in breast cancer and other solid tumors. Besides amplifications of known oncogenes, over

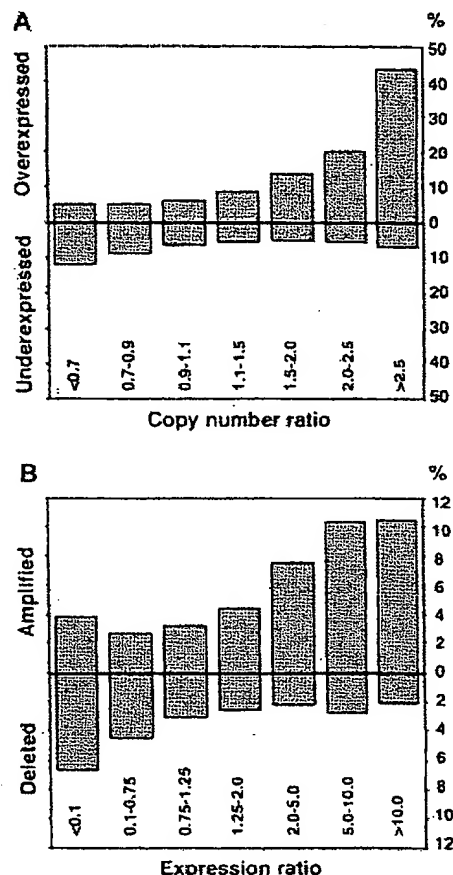


Fig. 1. Impact of gene copy number on global gene expression levels. A, percentage of over- and underexpressed genes (Y axis) according to copy number ratios (X axis). Threshold values used for over- and underexpression were >2.184 (global upper 7% of the cDNA ratios) and <0.4826 (global lower 7% of the expression ratios). B, percentage of amplified and deleted genes according to expression ratios. Threshold values for amplification and deletion were >1.5 and <0.7 .

20 recurrent regions of DNA amplification have been mapped in breast cancer by CGH⁵ (9, 10). However, these amplicons are often large and poorly defined, and their impact on gene expression remains unknown.

We hypothesized that genome-wide identification of those gene expression changes that are attributable to underlying gene copy number alterations would highlight transcripts that are actively involved in the causation or maintenance of the malignant phenotype. To identify such transcripts, we applied a combination of cDNA and CGH microarrays to: (a) determine the global impact that gene copy number variation plays in breast cancer development and progression; and (b) identify and characterize those genes whose mRNA expres-

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² Supplementary data for this article are available at Cancer Research Online (<http://cancerres.aacrjournals.org>).

³ Contributed equally to this work.

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⁵ The abbreviations used are: CGH, comparative genomic hybridization; FISH, fluorescence in situ hybridization; RT-PCR, reverse transcription-PCR.

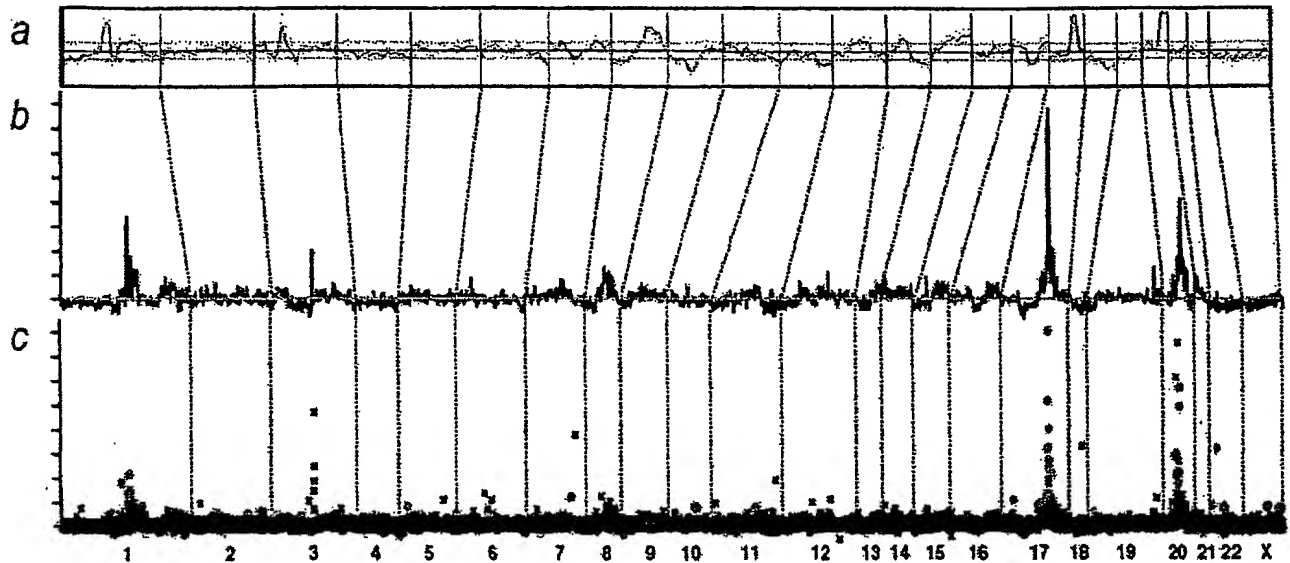


Fig. 2. Genome-wide copy number and expression analysis in the MCF-7 breast cancer cell line. *A*, chromosomal CGH analysis of MCF-7. The copy number ratio profile (blue line) across the entire genome from 1p telomere to Xq telomere is shown along with ± 1 SD (orange lines). The black horizontal line indicates a ratio of 1.0; red line, a ratio of 0.8; and green line, a ratio of 1.2. *B-C*, genome-wide copy number analysis in MCF-7 by CGH on cDNA microarray. The copy number ratios were plotted as a function of the position of the cDNA clones along the human genome. In *B*, individual data points are connected with a line, and a moving median of 10 adjacent clones is shown. Red horizontal line, the copy number ratio of 1.0. In *C*, individual data points are labeled by color coding according to cDNA expression ratios. The bright red dots indicate the upper 2%, and dark red dots, the next 5% of the expression ratios in MCF-7 cells (overexpressed genes); bright green dots indicate the lowest 2%, and dark green dots, the next 5% of the expression ratios (underexpressed genes); the rest of the observations are shown with black crosses. The chromosome numbers are shown at the bottom of the figure, and chromosome boundaries are indicated with a dashed line.

sion is most significantly associated with amplification of the corresponding genomic template.

MATERIALS AND METHODS

Breast Cancer Cell Lines. Fourteen breast cancer cell lines (BT-20, BT-474, HCC1428, Hs578t, MCF7, MDA-361, MDA-436, MDA-453, MDA-468, SKBR-3, T-47D, UACC812, ZR-75-1, and ZR-75-30) were obtained from the American Type Culture Collection (Manassas, VA). Cells were grown under recommended culture conditions. Genomic DNA and mRNA were isolated using standard protocols.

Copy Number and Expression Analyses by cDNA Microarrays. The preparation and printing of the 13,824 cDNA clones on glass slides were performed as described (11–13). Of these clones, 244 represented uncharacterized expressed sequence tags, and the remainder corresponded to known genes. CGH experiments on cDNA microarrays were done as described (14, 15). Briefly, 20 μ g of genomic DNA from breast cancer cell lines and normal human WBCs were digested for 14–18 h with *Ahl* and *RsaI* (Life Technologies, Inc., Rockville, MD) and purified by phenol/chloroform extraction. Six μ g of digested cell line DNAs were labeled with Cy3-dUTP (Amersham Pharmacia) and normal DNA with Cy5-dUTP (Amersham Pharmacia) using the Bioprime Labeling kit (Life Technologies, Inc.). Hybridization (14, 15) and posthybridization washes (13) were done as described. For the expression analyses, a standard reference (Universal Human Reference RNA; Stratagene, La Jolla, CA) was used in all experiments. Forty μ g of reference RNA were labeled with Cy3-dUTP and 3.5 μ g of test mRNA with Cy5-dUTP, and the labeled cDNAs were hybridized on microarrays as described (13, 15). For both microarray analyses, a laser confocal scanner (Agilent Technologies, Palo Alto, CA) was used to measure the fluorescence intensities at the target locations using the DEARRAY software (16). After background subtraction, average intensities at each clone in the test hybridization were divided by the average intensity of the corresponding clone in the control hybridization. For the copy number analysis, the ratios were normalized on the basis of the distribution of ratios of all targets on the array and for the expression analysis on the basis of 88 housekeeping genes, which were spotted four times onto the array. Low quality measurements (*i.e.*, copy number data with mean reference intensity <100 fluorescent units, and expression data with both test and reference intensity <100 fluorescent units and/or with spot size <50 units)

were excluded from the analysis and were treated as missing values. The distributions of fluorescence ratios were used to define cutpoints for increased/decreased copy number. Genes with CGH ratio >1.43 (representing the upper 5% of the CGH ratios across all experiments) were considered to be amplified, and genes with ratio <0.73 (representing the lower 5%) were considered to be deleted.

Statistical Analysis of CGH and cDNA Microarray Data. To evaluate the influence of copy number alterations on gene expression, we applied the following statistical approach. CGH and cDNA calibrated intensity ratios were log-transformed and normalized using median centering of the values in each cell line. Furthermore, cDNA ratios for each gene across all 14 cell lines were median centered. For each gene, the CGH data were represented by a vector that was labeled 1 for amplification (ratio, >1.43) and 0 for no amplification. Amplification was correlated with gene expression using the signal-to-noise statistics (1). We calculated a weight, w_g , for each gene as follows:

$$w_g = \frac{m_{g1} - m_{g0}}{\sigma_{g1} + \sigma_{g0}}$$

where m_{g1} , σ_{g1} and m_{g0} , σ_{g0} denote the means and SDs for the expression levels for amplified and nonamplified cell lines, respectively. To assess the statistical significance of each weight, we performed 10,000 random permutations of the label vector. The probability that a gene had a larger or equal weight by random permutation than the original weight was denoted by α . A low α (<0.05) indicates a strong association between gene expression and amplification.

Genomic Localization of cDNA Clones and Amplicon Mapping. Each cDNA clone on the microarray was assigned to a Unigene cluster using the Unigene Build 141.⁶ A database of genomic sequence alignment information for mRNA sequences was created from the August 2001 freeze of the University of California Santa Cruz's GoldenPath database.⁷ The chromosome and bp positions for each cDNA clone were then retrieved by relating these data sets. Amplicons were defined as a CGH copy number ratio >2.0 in at least two adjacent clones in two or more cell lines or a CGH ratio >2.0 in at least three adjacent clones in a single cell line. The amplicon start and end positions were

⁶ Internet address: http://research.nhgri.nih.gov/microarray/downloadable_cdna.html.

⁷ Internet address: www.genome.ucsc.edu.

Table 1 Summary of independent amplicons in 14 breast cancer cell lines by CGH microarray

Location	Start (Mb)	End (Mb)	Size (Mb)
1p13	132.79	132.94	0.2
1q21	173.92	177.25	3.3
1q22	179.28	179.57	0.3
3p14	71.94	74.66	2.7
7p12.1-7p11.2	55.62	60.95	5.3
7q31	125.73	130.96	5.2
7q32	140.01	140.68	0.7
8q21.11-8q21.13	86.45	92.46	6.0
8q21.3	98.45	103.05	4.6
8q23.3-8q24.14	129.88	142.15	12.3
8q24.22	151.21	152.16	1.0
9p13	38.65	39.25	0.6
13q22-q31	77.15	81.38	4.2
16q22	86.70	87.62	0.9
17q11	29.30	30.85	1.6
17q12-q21.2	39.79	42.80	3.0
17q21.32-q21.33	52.47	55.80	3.3
17q22-q23.3	63.81	69.70	5.9
17q23.3-q24.3	69.93	74.99	5.1
19q13	40.63	41.40	0.8
20q11.22	34.59	35.85	1.3
20q13.12	44.00	45.62	1.6
20q13.12-q13.13	46.45	49.43	3.0
20q13.2-q13.32	51.32	59.12	7.8

extended to include neighboring nonamplified clones (ratio, <1.5). The amplicon size determination was partially dependent on local clone density.

FISH. Dual-color interphase FISH to breast cancer cell lines was done as described (17). Bacterial artificial chromosome clone RP11-361K8 was labeled with SpectrumOrange (Vysis, Downers Grove, IL), and SpectrumOrange-labeled probe for *EGFR* was obtained from Vysis. SpectrumGreen-labeled chromosome 7 and 17 centromere probes (Vysis) were used as a reference. A tissue microarray containing 612 formalin-fixed, paraffin-embedded primary breast cancers (17) was applied in FISH analyses as described (18). The use of these specimens was approved by the Ethics Committee of the University of Basel and by the NIH. Specimens containing a 2-fold or higher increase in the number of test probe signals, as compared with corresponding centromere signals, in at least 10% of the tumor cells were considered to be amplified. Survival analysis was performed using the Kaplan-Meier method and the log-rank test.

RT-PCR. The *HOXB7* expression level was determined relative to *GAPDH*. Reverse transcription and PCR amplification were performed using Access RT-PCR System (Promega Corp., Madison, WI) with 10 ng of mRNA as a template. *HOXB7* primers were 5'-GAGCAGAGGGACTCGGACTT-3' and 5'-GCGTCAGGTAGCGATTGTAG-3'.

RESULTS

Global Effect of Copy Number on Gene Expression. 13,824 arrayed cDNA clones were applied for analysis of gene expression and gene copy number (CGH microarrays) in 14 breast cancer cell lines. The results illustrate a considerable influence of copy number on gene expression patterns. Up to 44% of the highly amplified transcripts (CGH ratio, >2.5) were overexpressed (i.e., belonged to the global upper 7% of expression ratios), compared with only 6% for genes with normal copy number levels (Fig. 1A). Conversely, 10.5% of the transcripts with high-level expression (cDNA ratio, >10) showed increased copy number (Fig. 1B). Low-level copy number increases and decreases were also associated with similar, although less dramatic, outcomes on gene expression (Fig. 1).

Identification of Distinct Breast Cancer Amplicons. Base-pair locations obtained for 11,994 cDNAs (86.8%) were used to plot copy number changes as a function of genomic position (Fig. 2, Supplement Fig. A). The average spacing of clones throughout the genome was 267 kb. This high-resolution mapping identified 24 independent breast cancer amplicons, spanning from 0.2 to 12 Mb of DNA (Table 1). Several amplification sites detected previously by chromosomal

CGH were validated, with 1q21, 17q12-q21.2, 17q22-q23, 20q13.1, and 20q13.2 regions being most commonly amplified. Furthermore, the boundaries of these amplicons were precisely delineated. In addition, novel amplicons were identified at 9p13 (38.65-39.25 Mb), and 17q21.3 (52.47-55.80 Mb).

Direct Identification of Putative Amplification Target Genes. The cDNA/CGH microarray technique enables the direct correlation of copy number and expression data on a gene-by-gene basis throughout the genome. We directly annotated high-resolution CGH plots with gene expression data using color coding. Fig. 2C shows that most of the amplified genes in the MCF-7 breast cancer cell line at 1p13, 17q22-q23, and 20q13 were highly overexpressed. A view of chromosome 7 in the MDA-468 cell line implicates *EGFR* as the most highly overexpressed and amplified gene at 7p11-p12 (Fig. 3A). In BT-474, the two known amplicons at 17q12 and 17q22-q23 contained numerous highly overexpressed genes (Fig. 3B). In addition, several genes, including the homeobox genes *HOXB2* and *HOXB7*, were highly amplified in a previously undescribed independent amplicon at 17q21.3. *HOXB7* was systematically amplified (as validated by FISH, Fig. 3B, inset) as well as overexpressed (as verified by RT-PCR, data not shown) in BT-474, UACC812, and ZR-75-30 cells. Furthermore, this novel

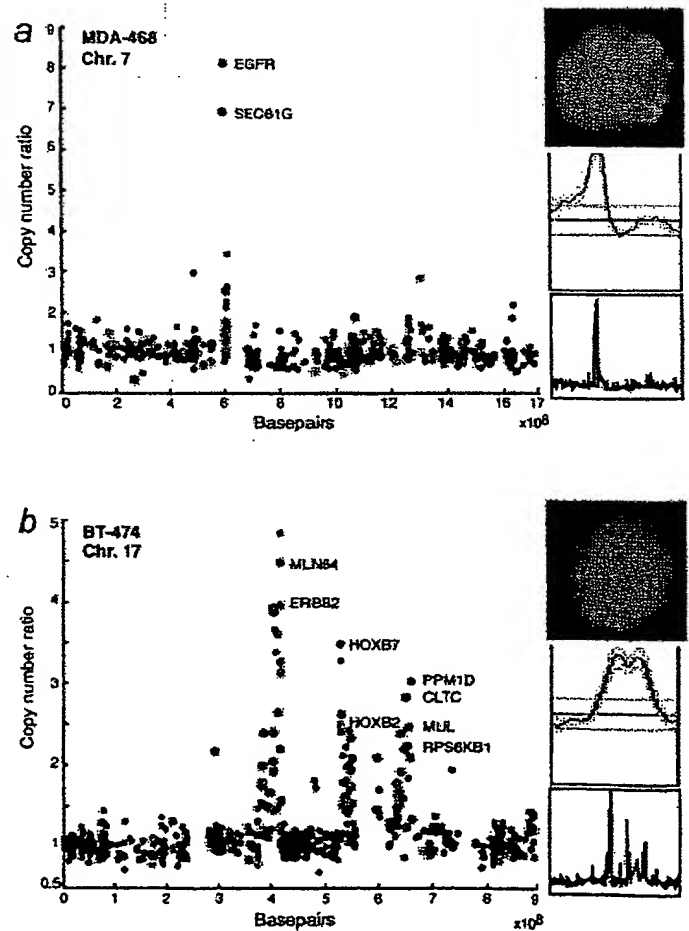
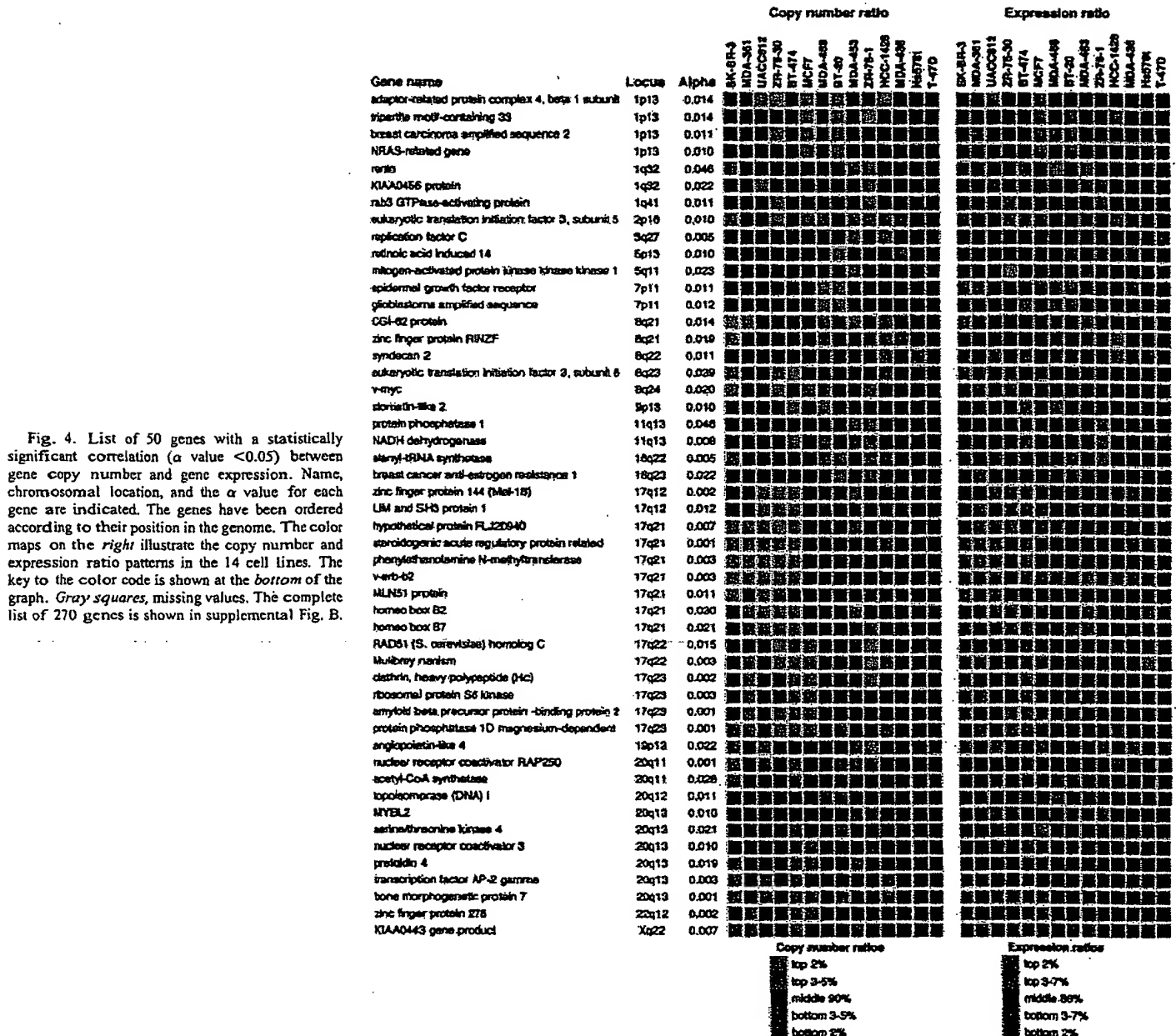


Fig. 3. Annotation of gene expression data on CGH microarray profiles. A, genes in the 7p11-p12 amplicon in the MDA-468 cell line are highly expressed (red dots) and include the *EGFR* oncogene. B, several genes in the 17q12, 17q21.3, and 17q23 amplicons in the BT-474 breast cancer cell line are highly overexpressed (red) and include the *HOXB7* gene. The data labels and color coding are as indicated for Fig. 2C. Insets show chromosomal CGH profiles for the corresponding chromosomes and validation of the increased copy number by interphase FISH using *EGFR* (red) and chromosome 7 centromere probe (green) to MDA-468 (A) and *HOXB7*-specific probe (red) and chromosome 17 centromere (green) to BT-474 cells (B).



amplification was validated to be present in 10.2% of 363 primary breast cancers by FISH to a tissue microarray and was associated with poor prognosis of the patients ($P = 0.001$).

Statistical Identification and Characterization of 270 Highly Expressed Genes in Amplicons. Statistical comparison of expression levels of all genes as a function of gene amplification identified 270 genes whose expression was significantly influenced by copy number across all 14 cell lines (Fig. 4, Supplemental Fig. B). According to the gene ontology data,⁸ 91 of the 270 genes represented hypothetical proteins or genes with no functional annotation, whereas 179 had associated functional information available. Of these, 151 (84%) are implicated in apoptosis, cell proliferation, signal transduction, and transcription, whereas 28 (16%) had functional annotations that could not be directly linked with cancer.

DISCUSSION

The importance of recurrent gene and chromosome copy number changes in the development and progression of solid tumors has been characterized in >1000 publications applying CGH⁹ (9, 10), as well as in a large number of other molecular cytogenetic, cytogenetic, and molecular genetic studies. The effects of these somatic genetic changes on gene expression levels have remained largely unknown, although a few studies have explored gene expression changes occurring in specific amplicons (15, 19–21). Here, we applied genome-wide cDNA microarrays to identify transcripts whose expression changes were attributable to underlying gene copy number alterations in breast cancer.

The overall impact of copy number on gene expression patterns was substantial with the most dramatic effects seen in the case of high-

⁸ Internet address: <http://www.geneontology.org/>.

⁹ Internet address: <http://www.ncbi.nlm.nih.gov/entrez>.

level copy number increase. Low-level copy number gains and losses also had a significant influence on expression levels of genes in the regions affected, but these effects were more subtle on a gene-by-gene basis than those of high-level amplifications. However, the impact of low-level gains on the dysregulation of gene expression patterns in cancer may be equally important if not more important than that of high-level amplifications. Aneuploidy and low-level gains and losses of chromosomal arms represent the most common types of genetic alterations in breast and other cancers and, therefore, have an influence on many genes. Our results in breast cancer extend the recent studies on the impact of aneuploidy on global gene expression patterns in yeast cells, acute myeloid leukemia, and a prostate cancer model system (22–24).

The CGH microarray analysis identified 24 independent breast cancer amplicons. We defined the precise boundaries for many amplicons detected previously by chromosomal CGH (9, 10, 25, 26) and also discovered novel amplicons that had not been detected previously, presumably because of their small size (only 1–2 Mb) or close proximity to other larger amplicons. One of these novel amplicons involved the homeobox gene region at 17q21.3 and led to the overexpression of the *HOXB7* and *HOXB2* genes. The homeodomain transcription factors are known to be key regulators of embryonic development and have been occasionally reported to undergo aberrant expression in cancer (27, 28). *HOXB7* transfection induced cell proliferation in melanoma, breast, and ovarian cancer cells and increased tumorigenicity and angiogenesis in breast cancer (29–32). The present results imply that gene amplification may be a prominent mechanism for overexpressing *HOXB7* in breast cancer and suggest that *HOXB7* contributes to tumor progression and confers an aggressive disease phenotype in breast cancer. This view is supported by our finding of amplification of *HOXB7* in 10% of 363 primary breast cancers, as well as an association of amplification with poor prognosis of the patients.

We carried out a systematic search to identify genes whose expression levels across all 14 cell lines were attributable to amplification status. Statistical analysis revealed 270 such genes (representing ~2% of all genes on the array), including not only previously described amplified genes, such as *HER-2*, *MYC*, *EGFR*, ribosomal protein *s6* kinase, and *AIB3*, but also numerous novel genes such as *NRAS-related gene* (1p13), *syndecan-2* (8q22), and *bone morphogenic protein* (20q13.1), whose activation by amplification may similarly promote breast cancer progression. Most of the 270 genes have not been implicated previously in breast cancer development and suggest novel pathogenetic mechanisms. Although we would not expect all of them to be causally involved, it is intriguing that 84% of the genes with associated functional information were implicated in apoptosis, cell proliferation, signal transduction, transcription, or other cellular processes that could directly imply a possible role in cancer progression. Therefore, a detailed characterization of these genes may provide biological insights to breast cancer progression and might lead to the development of novel therapeutic strategies.

In summary, we demonstrate application of cDNA microarrays to the analysis of both copy number and expression levels of over 12,000 transcripts throughout the breast cancer genome, roughly once every 267 kb. This analysis provided: (a) evidence of a prominent global influence of copy number changes on gene expression levels; (b) a high-resolution map of 24 independent amplicons in breast cancer; and (c) identification of a set of 270 genes, the overexpression of which was statistically attributable to gene amplification. Characterization of a novel amplicon at 17q21.3 implicated amplification and overexpression of the *HOXB7* gene in breast cancer, including a clinical association

between *HOXB7* amplification and poor patient prognosis. Overall, our results illustrate how the identification of genes activated by gene amplification provides a powerful approach to highlight genes with an important role in cancer as well as to prioritize and validate putative targets for therapy development.

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Microarray analysis reveals a major direct role of DNA copy number alteration in the transcriptional program of human breast tumors

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Genomic DNA copy number alterations are key genetic events in the development and progression of human cancers. Here we report a genome-wide microarray comparative genomic hybridization (array CGH) analysis of DNA copy number variation in a series of primary human breast tumors. We have profiled DNA copy number alteration across 6,691 mapped human genes, in 44 predominantly advanced, primary breast tumors and 10 breast cancer cell lines. While the overall patterns of DNA amplification and deletion corroborate previous cytogenetic studies, the high-resolution (gene-by-gene) mapping of amplicon boundaries and the quantitative analysis of amplicon shape provide significant improvement in the localization of candidate oncogenes. Parallel microarray measurements of mRNA levels reveal the remarkable degree to which variation in gene copy number contributes to variation in gene expression in tumor cells. Specifically, we find that 62% of highly amplified genes show moderately or highly elevated expression, that DNA copy number influences gene expression across a wide range of DNA copy number alterations (deletion, low-, mid- and high-level amplification), that on average, a 2-fold change in DNA copy number is associated with a corresponding 1.5-fold change in mRNA levels, and that overall, at least 12% of all the variation in gene expression among the breast tumors is directly attributable to underlying variation in gene copy number. These findings provide evidence that widespread DNA copy number alteration can lead directly to global deregulation of gene expression, which may contribute to the development or progression of cancer.

Conventional cytogenetic techniques, including comparative genomic hybridization (CGH) (1), have led to the identification of a number of recurrent regions of DNA copy number alteration in breast cancer cell lines and tumors (2–4). While some of these regions contain known or candidate oncogenes [e.g., *FGFR1* (8p11), *MYC* (8q24), *CCND1* (11q13), *ERBB2* (17q12), and *ZNF217* (20q13)] and tumor suppressor genes [*RB1* (13q14) and *TP53* (17p13)], the relevant gene(s) within other regions (e.g., gain of 1q, 8q22, and 17q22–24, and loss of 8p) remain to be identified. A high-resolution genome-wide map, delineating the boundaries of DNA copy number alterations in tumors, should facilitate the localization and identification of oncogenes and tumor suppressor genes in breast cancer. In this study, we have created such a map, using array-based CGH (5–7) to profile DNA copy number alteration in a series of breast cancer cell lines and primary tumors.

An unresolved question is the extent to which the widespread DNA copy number changes that we and others have identified in breast tumors alter expression of genes within involved regions. Because we had measured mRNA levels in parallel in the same samples (8), using the same DNA microarrays, we had an opportunity to explore on a genomic scale the relationship between DNA copy number changes and gene expression. From

this analysis, we have identified a significant impact of widespread DNA copy number alteration on the transcriptional programs of breast tumors.

Materials and Methods

Tumors and Cell Lines. Primary breast tumors were predominantly large (>3 cm), intermediate-grade, infiltrating ductal carcinomas, with more than 50% being lymph node positive. The fraction of tumor cells within specimens averaged at least 50%. Details of individual tumors have been published (8, 9), and are summarized in Table 1, which is published as supporting information on the PNAS web site, www.pnas.org. Breast cancer cell lines were obtained from the American Type Culture Collection. Genomic DNA was isolated either using Qiagen genomic DNA columns, or by phenol/chloroform extraction followed by ethanol precipitation.

DNA Labeling and Microarray Hybridizations. Genomic DNA labeling and hybridizations were performed essentially as described in Pollack *et al.* (7), with slight modifications. Two micrograms of DNA was labeled in a total volume of 50 microliters and the volumes of all reagents were adjusted accordingly. “Test” DNA (from tumors and cell lines) was fluorescently labeled (Cy5) and hybridized to a human cDNA microarray containing 6,691 different mapped human genes (i.e., UniGene clusters). The “reference” (labeled with Cy3) for each hybridization was normal female leukocyte DNA from a single donor. The fabrication of cDNA microarrays and the labeling and hybridization of mRNA samples have been described (8).

Data Analysis and Map Positions. Hybridized arrays were scanned on a GenePix scanner (Axon Instruments, Foster City, CA), and fluorescence ratios (test/reference) calculated using SCANALYZE software (available at <http://rana.lbl.gov>). Fluorescence ratios were normalized for each array by setting the average log fluorescence ratio for all array elements equal to 0. Measurements with fluorescence intensities more than 20% above background were considered reliable. DNA copy number profiles that deviated significantly from background ratios measured in normal genomic DNA control hybridizations were interpreted as evidence of real DNA copy number alteration (see *Estimating Significance of Altered Fluorescence Ratios* in the supporting information). When indicated, DNA copy number profiles are displayed as a moving average (symmetric 5-nearest neighbors). Map positions for arrayed human cDNAs were assigned by

Abbreviation: CGH, comparative genomic hybridization.

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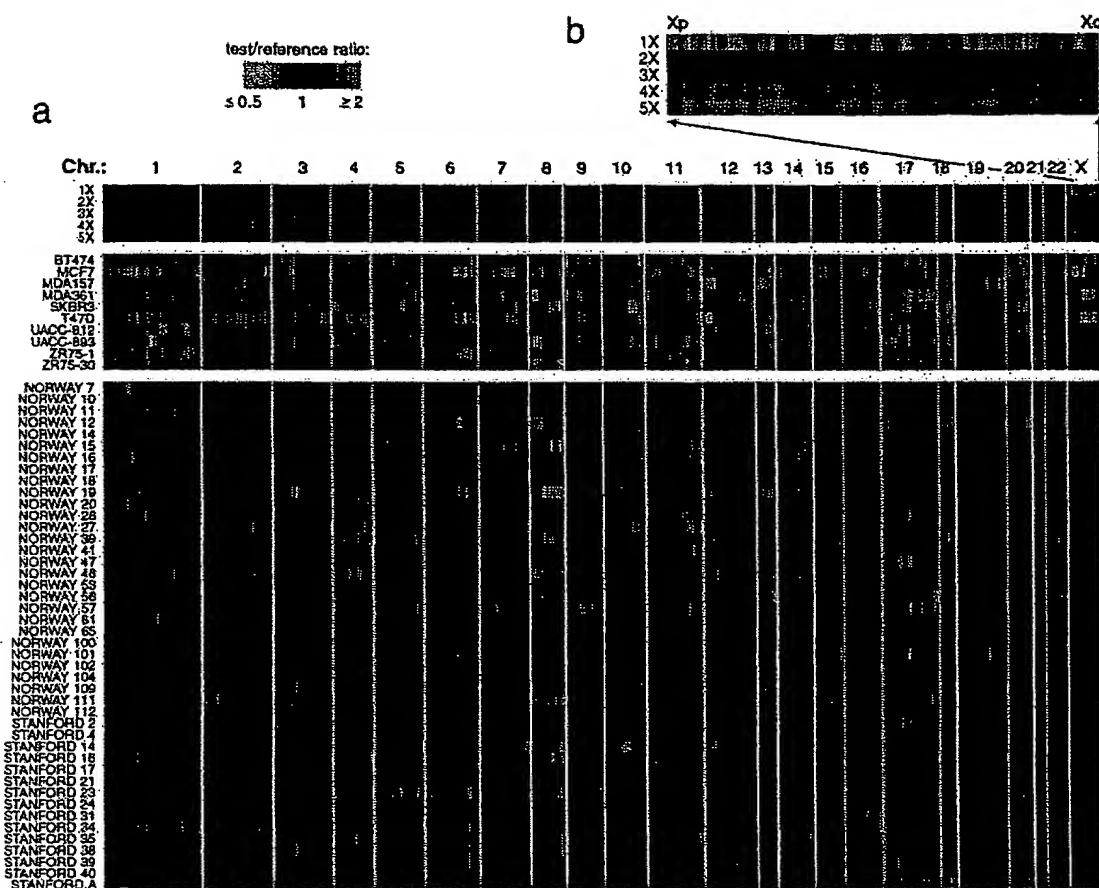


Fig. 1. Genome-wide measurement of DNA copy number alteration by array CGH. (a) DNA copy number profiles are illustrated for cell lines containing different numbers of X chromosomes, for breast cancer cell lines, and for breast tumors. Each row represents a different cell line or tumor, and each column represents one of 6,691 different mapped human genes present on the microarray, ordered by genome map position from 1pter through Xqter. Moving average (symmetric 5-nearest neighbors) fluorescence ratios (test/reference) are depicted using a \log_2 -based pseudocolor scale (indicated), such that red luminescence reflects fold-amplification, green luminescence reflects fold-deletion, and black indicates no change (gray indicates poorly measured data). (b) Enlarged view of DNA copy number profiles across the X chromosome, shown for cell lines containing different numbers of X chromosomes.

identifying the starting position of the best and longest match of any DNA sequence represented in the corresponding UniGene cluster (10) against the "Golden Path" genome assembly (<http://genome.ucsc.edu/>; Oct 7, 2000 Freeze). For UniGene clusters represented by multiple arrayed elements, mean fluorescence ratios (for all elements representing the same UniGene cluster) are reported. For mRNA measurements, fluorescence ratios are "mean-centered" (i.e., reported relative to the mean ratio across the 44 tumor samples). The data set described here can be accessed in its entirety in the supporting information.

Results

We performed CGH on 44 predominantly locally advanced, primary breast tumors and 10 breast cancer cell lines, using cDNA microarrays containing 6,691 different mapped human genes (Fig. 1a; also see *Materials and Methods* for details of microarray hybridizations). To take full advantage of the improved spatial resolution of array CGH, we ordered (fluorescence ratios for) the 6,691 cDNAs according to the "Golden Path" (<http://genome.ucsc.edu/>) genome assembly of the draft human genome sequences (11). In so doing, arrayed cDNAs not only themselves represent genes of potential interest (e.g., candidate oncogenes within amplicons), but also provide precise genetic landmarks for chromosomal regions of amplification and

deletion. Parallel analysis of DNA from cell lines containing different numbers of X chromosomes (Fig. 1b), as we did before (7), demonstrated the sensitivity of our method to detect single-copy loss (45, XO), and 1.5- (47, XXX), 2- (48, XXXX), or 2.5-fold (49, XXXXX) gains (also see Fig. 5, which is published as supporting information on the PNAS web site). Fluorescence ratios were linearly proportional to copy number ratios, which were slightly underestimated, in agreement with previous observations (7). Numerous DNA copy number alterations were evident in both the breast cancer cell lines and primary tumors (Fig. 1a), detected in the tumors despite the presence of euploid non-tumor cell types; the magnitudes of the observed changes were generally lower in the tumor samples. DNA copy-number alterations were found in every cancer cell line and tumor, and on every human chromosome in at least one sample. Recurrent regions of DNA copy number gain and loss were readily identifiable. For example, gains within 1q, 8q, 17q, and 20q were observed in a high proportion of breast cancer cell lines/tumors (90%/69%, 100%/47%, 100%/60%, and 90%/44%, respectively), as were losses within 1p, 3p, 8p, and 13q (80%/24%, 80%/22%, 80%/22%, and 70%/18%, respectively), consistent with published cytogenetic studies (refs. 2-4; a complete listing of gains/losses is provided in Tables 2 and 3, which are published as supporting information on the PNAS web site). The total

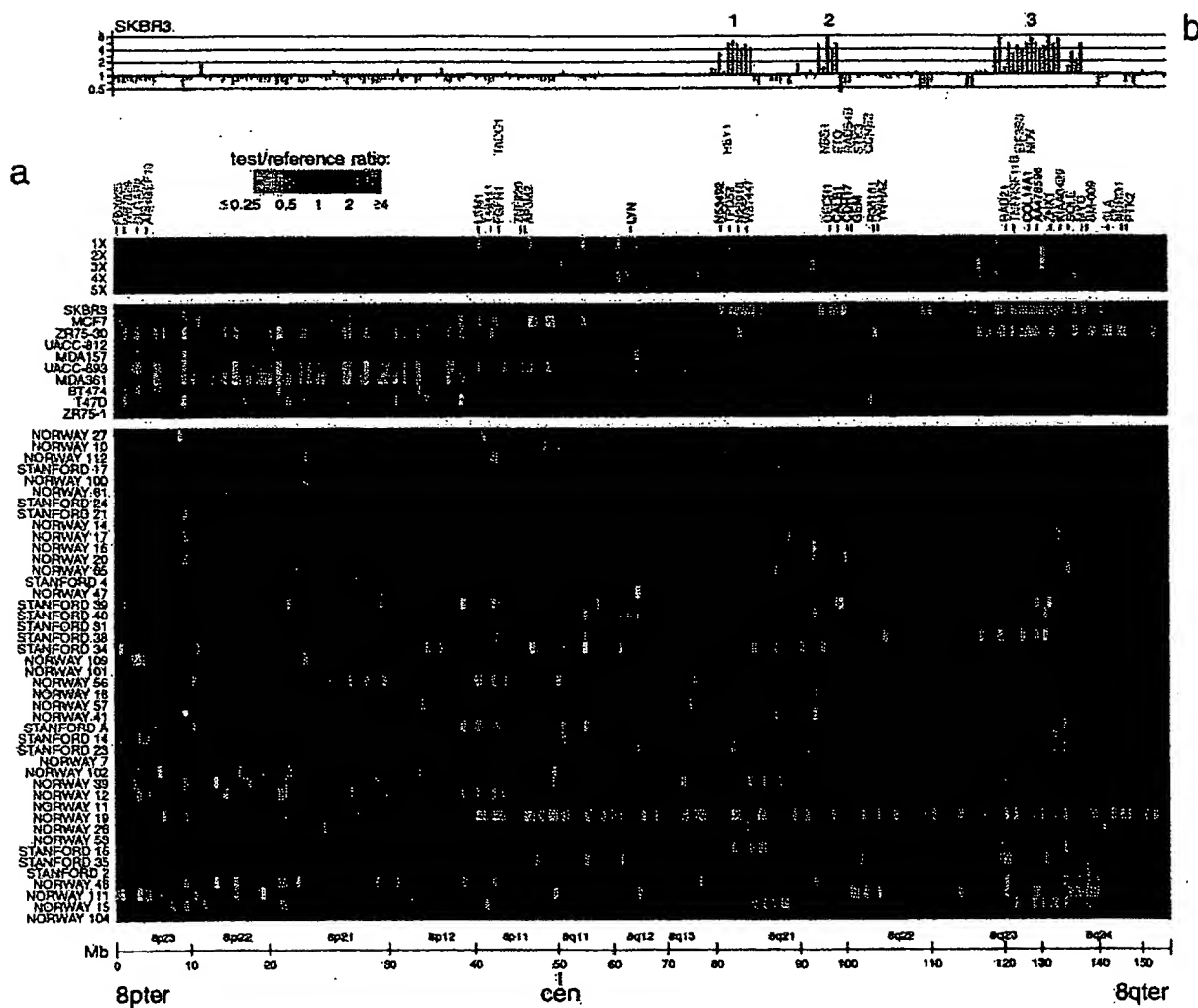


Fig. 2. DNA copy number alteration across chromosome 8 by array CGH. (a) DNA copy number profiles are illustrated for cell lines containing different numbers of X chromosomes, for breast cancer cell lines, and for breast tumors. Breast cancer cell lines and tumors are separately ordered by hierarchical clustering to highlight recurrent copy number changes. The 241 genes present on the microarrays and mapping to chromosome 8 are ordered by position along the chromosome. Fluorescence ratios (test/reference) are depicted by a log₂ pseudocolor scale (indicated). Selected genes are indicated with color-coded text (red, increased; green, decreased; black, no change; gray, not well measured) to reflect correspondingly altered mRNA levels (observed in the majority of the subset of samples displaying the DNA copy number change). The map positions for genes of interest that are not represented on the microarray are indicated in the row above those genes represented on the array. (b) Graphical display of DNA copy number profile for breast cancer cell line SKBR3. Fluorescence ratios (tumor/normal) are plotted on a log₂ scale for chromosome 8 genes, ordered along the chromosome.

number of genomic alterations (gains and losses) was found to be significantly higher in breast tumors that were high grade ($P = 0.008$), consistent with published CGH data (3), estrogen receptor negative ($P = 0.04$), and harboring TP53 mutations ($P = 0.0006$) (see Table 4, which is published as supporting information on the PNAS web site).

The improved spatial resolution of our array CGH analysis is illustrated for chromosome 8, which displayed extensive DNA copy number alteration in our series. A detailed view of the variation in the copy number of 241 genes mapping to chromosome 8 revealed multiple regions of recurrent amplification; each of these potentially harbors a different known or previously uncharacterized oncogene (Fig. 2a). The complexity of amplicon structure is most easily appreciated in the breast cancer cell line SKBR3. Although a conventional CGH analysis of 8q in SKBR3 identified only two distinct regions of amplification (12), we observed three distinct regions of high-level amplification (labeled 1–3 in Fig. 2b). For each of these regions we can define the

boundaries of the interval recurrently amplified in the tumors we examined; in each case, known or plausible candidate oncogenes can be identified (a description of these regions, as well as the recurrently amplified regions on chromosomes 17 and 20, can be found in Figs. 6 and 7, which are published as supporting information on the PNAS web site).

For a subset of breast cancer cell lines and tumors (4 and 37, respectively), and a subset of arrayed genes (6,095), mRNA levels were quantitatively measured in parallel by using cDNA microarrays (8). The parallel assessment of mRNA levels is useful in the interpretation of DNA copy number changes. For example, the highly amplified genes that are also highly expressed are the strongest candidate oncogenes within an amplicon. Perhaps more significantly, our parallel analysis of DNA copy number changes and mRNA levels provides us the opportunity to assess the global impact of widespread DNA copy number alteration on gene expression in tumor cells.

A strong influence of DNA copy number on gene expression is evident in an examination of the pseudocolor representations

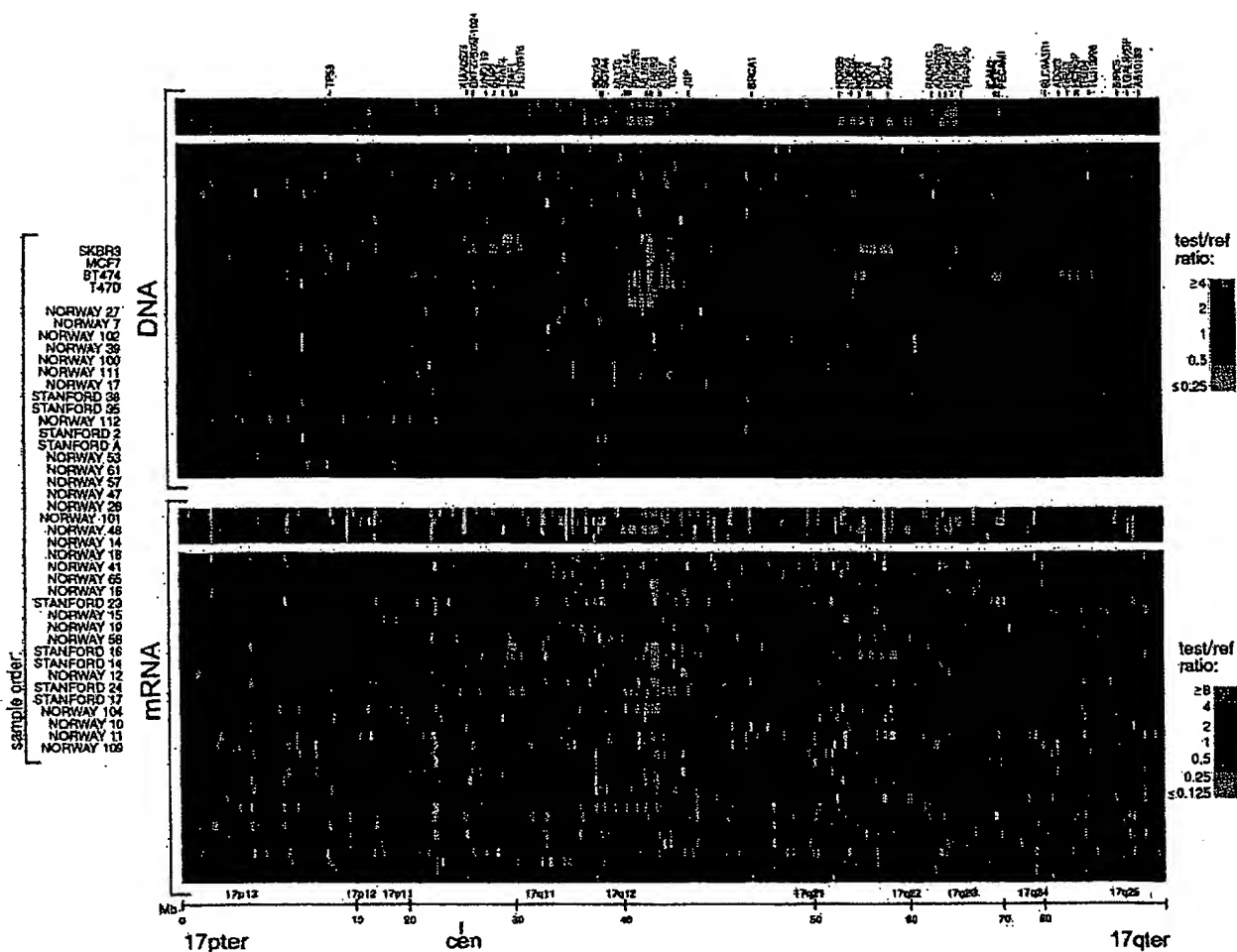


Fig. 3. Concordance between DNA copy number and gene expression across chromosome 17. DNA copy number alteration (Upper) and mRNA levels (Lower) are illustrated for breast cancer cell lines and tumors. Breast cancer cell lines and tumors are separately ordered by hierarchical clustering (Upper), and the identical sample order is maintained (Lower). The 354 genes present on the microarrays and mapping to chromosome 17, and for which both DNA copy number and mRNA levels were determined, are ordered by position along the chromosome; selected genes are indicated in color-coded text (see Fig. 2 legend). Fluorescence ratios (test/reference) are depicted by separate \log_2 pseudocolor scales (indicated).

of DNA copy number and mRNA levels for genes on chromosome 17 (Fig. 3). The overall patterns of gene amplification and elevated gene expression are quite concordant; i.e., a significant fraction of highly amplified genes appear to be correspondingly highly expressed. The concordance between high-level amplification and increased gene expression is not restricted to chromosome 17. Genome-wide, of 117 high-level DNA amplifications (fluorescence ratios >4 , and representing 91 different genes), 62% (representing 54 different genes; see Table 5, which is published as supporting information on the PNAS web site) are found associated with at least moderately elevated mRNA levels (mean-centered fluorescence ratios >2), and 42% (representing 36 different genes) are found associated with comparably highly elevated mRNA levels (mean-centered fluorescence ratios >4).

To determine the extent to which DNA deletion and lower-level amplification (in addition to high-level amplification) are also associated with corresponding alterations in mRNA levels, we performed three separate analyses on the complete data set (4 cell lines and 37 tumors, across 6,095 genes). First, we determined the average mRNA levels for each of five classes of genes, representing DNA deletion, no change, and low-, medium-, and high-level amplification (Fig. 4a). For both the

breast cancer cell lines and tumors, average mRNA levels tracked with DNA copy number across all five classes, in a statistically significant fashion (P values for pair-wise Student's t tests comparing adjacent classes: cell lines, 4×10^{-49} , 1×10^{-49} , 5×10^{-3} , 1×10^{-2} ; tumors, 1×10^{-43} , 1×10^{-214} , 5×10^{-41} , 1×10^{-4}). A linear regression of the average $\log(\text{DNA copy number})$, for each class, against average $\log(\text{mRNA level})$ demonstrated that on average, a 2-fold change in DNA copy number was accompanied by 1.4- and 1.5-fold changes in mRNA level for the breast cancer cell lines and tumors, respectively (Fig. 4a, regression line not shown). Second, we characterized the distribution of the 6,095 correlations between DNA copy number and mRNA level, each across the 37 tumor samples (Fig. 4b). The distribution of correlations forms a normal-shaped curve, but with the peak markedly shifted in the positive direction from zero. This shift is statistically significant, as evidenced in a plot of observed vs. expected correlations (Fig. 4c), and reflects a pervasive global influence of DNA copy number alterations on gene expression. Notably, the highest correlations between DNA copy number and mRNA level (the right tail of the distribution in Fig. 4b) comprise both amplified and deleted genes (data not shown). Third, we used a linear regression model to estimate the fraction of all variation measured in mRNA levels among the 37

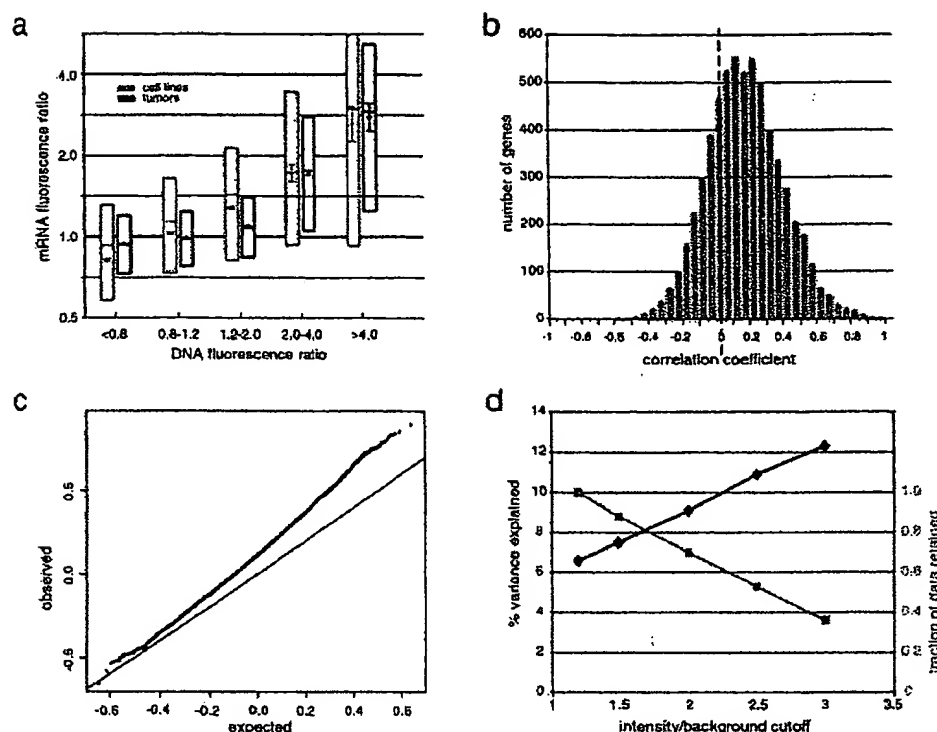


Fig. 4. Genome-wide influence of DNA copy number alterations on mRNA levels. (a) For breast cancer cell lines (gray) and tumor samples (black), both mean-centered mRNA fluorescence ratio (log₂ scale) quartiles (box plots indicate 25th, 50th, and 75th percentile) and averages (diamonds; Y-value error bars indicate standard errors of the mean) are plotted for each of five classes of genes, representing DNA deletion (tumor/normal ratio <0.8), no change (0.8–1.2), low- (1.2–2), medium- (2–4), and high-level (>4) amplification. *P* values for pair-wise Student's *t* tests, comparing averages between adjacent classes (moving left to right), are 4×10^{-69} , 1×10^{-49} , 5×10^{-5} , 1×10^{-2} (cell lines), and 1×10^{-43} , 1×10^{-214} , 5×10^{-41} , 1×10^{-4} (tumors). (b) Distribution of correlations between DNA copy number and mRNA levels, for 6,095 different human genes across 37 breast tumor samples. (c) Plot of observed versus expected correlation coefficients. The expected values were obtained by randomization of the sample labels in the DNA copy number data set. The line of unity is indicated. (d) Percent variance in gene expression (among tumors) directly explained by variation in gene copy number. Percent variance explained (black line) and fraction of data retained (gray line) are plotted for different fluorescence intensity/background (a rough surrogate for signal/noise) cutoff values. Fraction of data retained is relative to the 1.2 intensity/background cutoff. Details of the linear regression model used to estimate the fraction of variation in gene expression attributable to underlying DNA copy number alteration can be found in the supporting information (see *Estimating the Fraction of Variation in Gene Expression Attributable to Underlying DNA Copy Number Alteration*).

tumors that could be attributed to underlying variation in DNA copy number. From this analysis, we estimate that, overall, about 7% of all of the observed variation in mRNA levels can be explained directly by variation in copy number of the altered genes (Fig. 4d). We can reduce the effects of experimental measurement error on this estimate by using only that fraction of the data most reliably measured (fluorescence intensity/background >3); using that data, our estimate of the percent variation in mRNA levels directly attributed to variation in gene copy number increases to 12% (Fig. 4d). This still undoubtedly represents a significant underestimate, as the observed variation in global gene expression is affected not only by true variation in the expression programs of the tumor cells themselves, but also by the variable presence of non-tumor cell types within clinical samples.

Discussion

This genome-wide, array CGH analysis of DNA copy number alteration in a series of human breast tumors demonstrates the usefulness of defining amplicon boundaries at high resolution (gene-by-gene), and quantitatively measuring amplicon shape, to assist in locating and identifying candidate oncogenes. By analyzing mRNA levels in parallel, we have also discovered that changes in DNA copy number have a large, pervasive, direct effect on global gene expression patterns in both breast cancer

cell lines and tumors. Although the DNA microarrays used in our analysis may display a bias toward characterized and/or highly expressed genes, because we are examining such a large fraction of the genome (approximately 20% of all human genes), and because, as detailed above, we are likely underestimating the contribution of DNA copy number changes to altered gene expression, we believe our findings are likely to be generalizable (but would nevertheless still be remarkable if only applicable to this set of ~6,100 genes).

In budding yeast, aneuploidy has been shown to result in chromosome-wide gene expression biases (13). Two recent studies have begun to examine the global relationship between DNA copy number and gene expression in cancer cells. In agreement with our findings, Phillips *et al.* (14) have shown that with the acquisition of tumorigenicity in an immortalized prostate epithelial cell line, new chromosomal gains and losses resulted in a statistically significant respective increase and decrease in the average expression level of involved genes. In contrast, Platzer *et al.* (15) recently reported that in metastatic colon tumors only ~4% of genes within amplified regions were found more highly (>2-fold) expressed, when compared with normal colonic epithelium. This report differs substantially from our finding that 62% of highly amplified genes in breast cancer exhibit at least 2-fold increased expression. These contrasting findings may reflect methodological differences between the

studies. For example, the study of Platzer *et al.* (15) may have systematically under-measured gene expression changes. In this regard it is remarkable that only 14 transcripts of many thousand residing within unamplified chromosomal regions were found to exhibit at least 4-fold altered expression in metastatic colon cancer. Additionally, their reliance on lower-resolution chromosomal CGH may have resulted in poorly delimiting the boundaries of high-complexity amplicons, effectively overcalling regions with amplification. Alternatively, the contrasting findings for amplified genes may represent real biological differences between breast and metastatic colon tumors; resolution of this issue will require further studies.

Our finding that widespread DNA copy number alteration has a large, pervasive and direct effect on global gene expression patterns in breast cancer has several important implications. First, this finding supports a high degree of copy number-dependent gene expression in tumors. Second, it suggests that most genes are not subject to specific autoregulation or dosage compensation. Third, this finding cautions that elevated expression of an amplified gene cannot alone be considered strong independent evidence of a candidate oncogene's role in tumorigenesis. In our study, fully 62% of highly amplified genes demonstrated moderately or highly elevated expression. This highlights the importance of high-resolution mapping of amplicon boundaries and shape [to identify the "driving" gene(s) within amplicons (16)], on a large number of samples, in addition to functional studies. Fourth, this finding suggests that analyzing

the genomic distribution of expressed genes, even within existing microarray gene expression data sets, may permit the inference of DNA copy number aberration, particularly aneuploidy (where gene expression can be averaged across large chromosomal regions; see Fig. 3 and supporting information). Fifth, this finding implies that a substantial portion of the phenotypic uniqueness (and by extension, the heterogeneity in clinical behavior) among patients' tumors may be traceable to underlying variation in DNA copy number. Sixth, this finding supports a possible role for widespread DNA copy number alteration in tumorigenesis (17, 18), beyond the amplification of specific oncogenes and deletion of specific tumor suppressor genes. Widespread DNA copy number alteration, and the concomitant widespread imbalance in gene expression, might disrupt critical stoichiometric relationships in cell metabolism and physiology (e.g., proteasome, mitotic spindle), possibly promoting further chromosomal instability and directly contributing to tumor development or progression. Finally, our findings suggest the possibility of cancer therapies that exploit specific or global imbalances in gene expression in cancer.

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Genome-wide Study of Gene Copy Numbers, Transcripts, and Protein Levels in Pairs of Non-invasive and Invasive Human Transitional Cell Carcinomas*

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Gain and loss of chromosomal material is characteristic of bladder cancer, as well as malignant transformation in general. The consequences of these changes at both the transcription and translation levels is at present unknown partly because of technical limitations. Here we have attempted to address this question in pairs of non-invasive and invasive human bladder tumors using a combination of technology that included comparative genomic hybridization, high density oligonucleotide array-based monitoring of transcript levels (5600 genes), and high resolution two-dimensional gel electrophoresis. The results showed that there is a gene dosage effect that in some cases superimposes on other regulatory mechanisms. This effect depended ($p < 0.015$) on the magnitude of the comparative genomic hybridization change. In general (18 of 23 cases), chromosomal areas with more than 2-fold gain of DNA showed a corresponding increase in mRNA transcripts. Areas with loss of DNA, on the other hand, showed either reduced or unaltered transcript levels. Because most proteins resolved by two-dimensional gels are unknown it was only possible to compare mRNA and protein alterations in relatively few cases of well focused abundant proteins. With few exceptions we found a good correlation ($p < 0.005$) between transcript alterations and protein levels. The implications, as well as limitations, of the approach are discussed. *Molecular & Cellular Proteomics* 1:37–45, 2002.

Aneuploidy is a common feature of most human cancers (1), but little is known about the genome-wide effect of this

phenomenon at both the transcription and translation levels. High throughput array studies of the breast cancer cell line BT474 has suggested that there is a correlation between DNA copy numbers and gene expression in highly amplified areas (2), and studies of individual genes in solid tumors have revealed a good correlation between gene dose and mRNA or protein levels in the case of *c-erb-B2*, *cyclin d1*, *ems1*, and *N-myc* (3–5). However, a high cyclin D1 protein expression has been observed without simultaneous amplification (4), and a low level of *c-myc* copy number increase was observed without concomitant *c-myc* protein overexpression (6).

In human bladder tumors, karyotyping, fluorescent *in situ* hybridization, and comparative genomic hybridization (CGH)¹ have revealed chromosomal aberrations that seem to be characteristic of certain stages of disease progression. In the case of non-invasive pTa transitional cell carcinomas (TCCs), this includes loss of chromosome 9 or parts of it, as well as loss of Y in males. In minimally invasive pT1 TCCs, the following alterations have been reported: 2q–, 11p–, 1q+, 11q13+, 17q+, and 20q+ (7–12). It has been suggested that these regions harbor tumor suppressor genes and oncogenes; however, the large chromosomal areas involved often contain many genes, making meaningful predictions of the functional consequences of losses and gains very difficult.

In this investigation we have combined genome-wide technology for detecting genomic gains and losses (CGH) with gene expression profiling techniques (microarrays and proteomics) to determine the effect of gene copy number on transcript and protein levels in pairs of non-invasive and invasive human bladder TCCs.

EXPERIMENTAL PROCEDURES

Material—Bladder tumor biopsies were sampled after informed consent was obtained and after removal of tissue for routine pathology examination. By light microscopy tumors 335 and 532 were staged by an experienced pathologist as pTa (superficial papillary),

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¹ The abbreviations used are: CGH, comparative genomic hybridization; TCC, transitional cell carcinoma; LOH, loss of heterozygosity; PA-FABP, psoriasis-associated fatty acid-binding protein; 2D, two-dimensional.

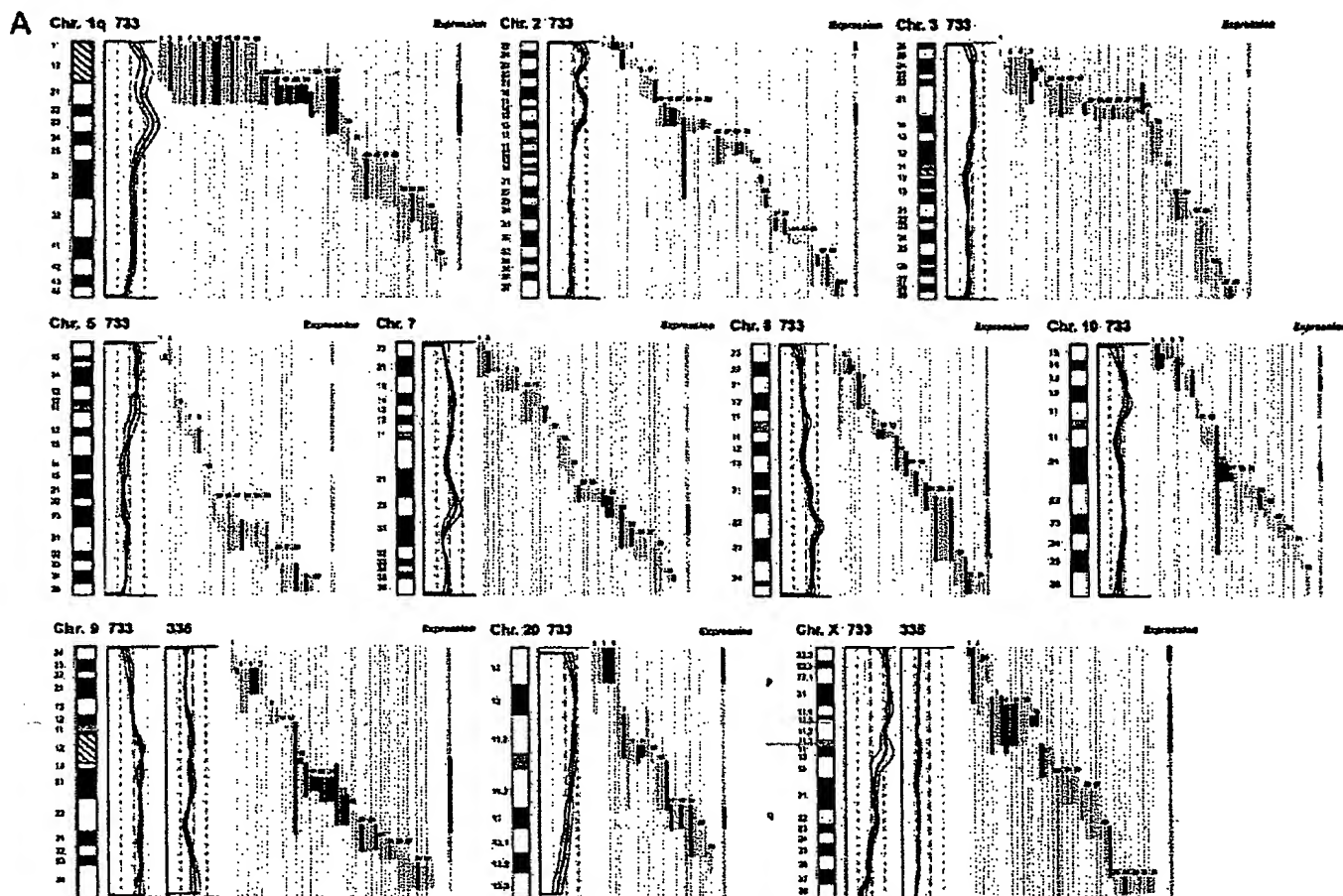


FIG. 1. DNA copy number and mRNA expression level. Shown from left to right are chromosome (Chr.), CGH profiles, gene location and expression level of specific genes, and overall expression level along the chromosome. A, expression of mRNA in invasive tumor 733 as compared with the non-invasive counterpart tumor 335. B, expression of mRNA in invasive tumor 827 compared with the non-invasive counterpart tumor 532. The average fluorescent signal ratio between tumor DNA and normal DNA is shown along the length of the chromosome (left). The bold curve in the ratio profile represents a mean of four chromosomes and is surrounded by thin curves indicating one standard deviation. The central vertical line (broken) indicates a ratio value of 1 (no change), and the vertical lines next to it (dotted) indicate a ratio of 0.5 (left) and 2.0 (right). In chromosomes where the non-invasive tumor 335 used for comparison showed alterations in DNA content, the ratio profile of that chromosome is shown to the right of the invasive tumor profile. The colored bars represent one gene each, identified by the running numbers above the bars (the name of the gene can be seen at www.MDL.DK/sdata.html). The bars indicate the purported location of the gene, and the colors indicate the expression level of the gene in the invasive tumor compared with the non-invasive counterpart; >2-fold increase (black), >2-fold decrease (blue), no significant change (orange). The bar to the far right, entitled Expression shows the resulting change in expression along the chromosome; the colors indicate that at least half of the genes were up-regulated (black), at least half of the genes down-regulated (blue), or more than half of the genes are unchanged (orange). If a gene was absent in one of the samples and present in another, it was regarded as more than a 2-fold change. A 2-fold level was chosen as this corresponded to one standard deviation in a double determination of ~1800 genes. Centromeres and heterochromatic regions were excluded from data analysis.

grade I and II, respectively, tumors 733 and 827 were staged as pT1 (invasive into submucosa), 733 was staged as solid, and 827 was staged as papillary, both grade III.

mRNA Preparation—Tissue biopsies, obtained fresh from surgery, were embedded immediately in a sodium-guanidinium thiocyanate solution and stored at -80°C . Total RNA was isolated using the RNeasy B RNA isolation method (WAK-Chemie Medical GmbH). poly(A)⁺ RNA was isolated by an oligo(dT) selection step (Oligotex mRNA kit; Qiagen).

cRNA Preparation—1 μg of mRNA was used as starting material. The first and second strand cDNA synthesis was performed using the SuperScript[®] choice system (Invitrogen) according to the manufacturer's instructions but using an oligo(dT) primer containing a T7 RNA polymerase binding site. Labeled cRNA was prepared using the MEGAscript[®] *in vitro* transcription kit (Ambion). Biotin-labeled CTP and

UTP (Enzo) was used, together with unlabeled NTPs in the reaction. Following the *in vitro* transcription reaction, the unincorporated nucleotides were removed using RNeasy columns (Qiagen).

Array Hybridization and Scanning—Array hybridization and scanning was modified from a previous method (13). 10 μg of cRNA was fragmented at 94°C for 35 min in buffer containing 40 mM Tris acetate, pH 8.1, 100 mM KOAc, 30 mM MgOAc. Prior to hybridization, the fragmented cRNA in a 6 \times SSPE-T hybridization buffer (1 M NaCl, 10 mM Tris, pH 7.6, 0.005% Triton), was heated to 95°C for 5 min, subsequently cooled to 40°C , and loaded onto the Affymetrix probe array cartridge. The probe array was then incubated for 16 h at 40°C at constant rotation (60 rpm). The probe array was exposed to 10 washes in 6 \times SSPE-T at 25°C followed by 4 washes in 0.5 \times SSPE-T at 50°C . The biotinylated cRNA was stained with a streptavidin-phycoerythrin conjugate, 10 $\mu\text{g}/\text{ml}$ (Molecular Probes) in 6 \times SSPE-T

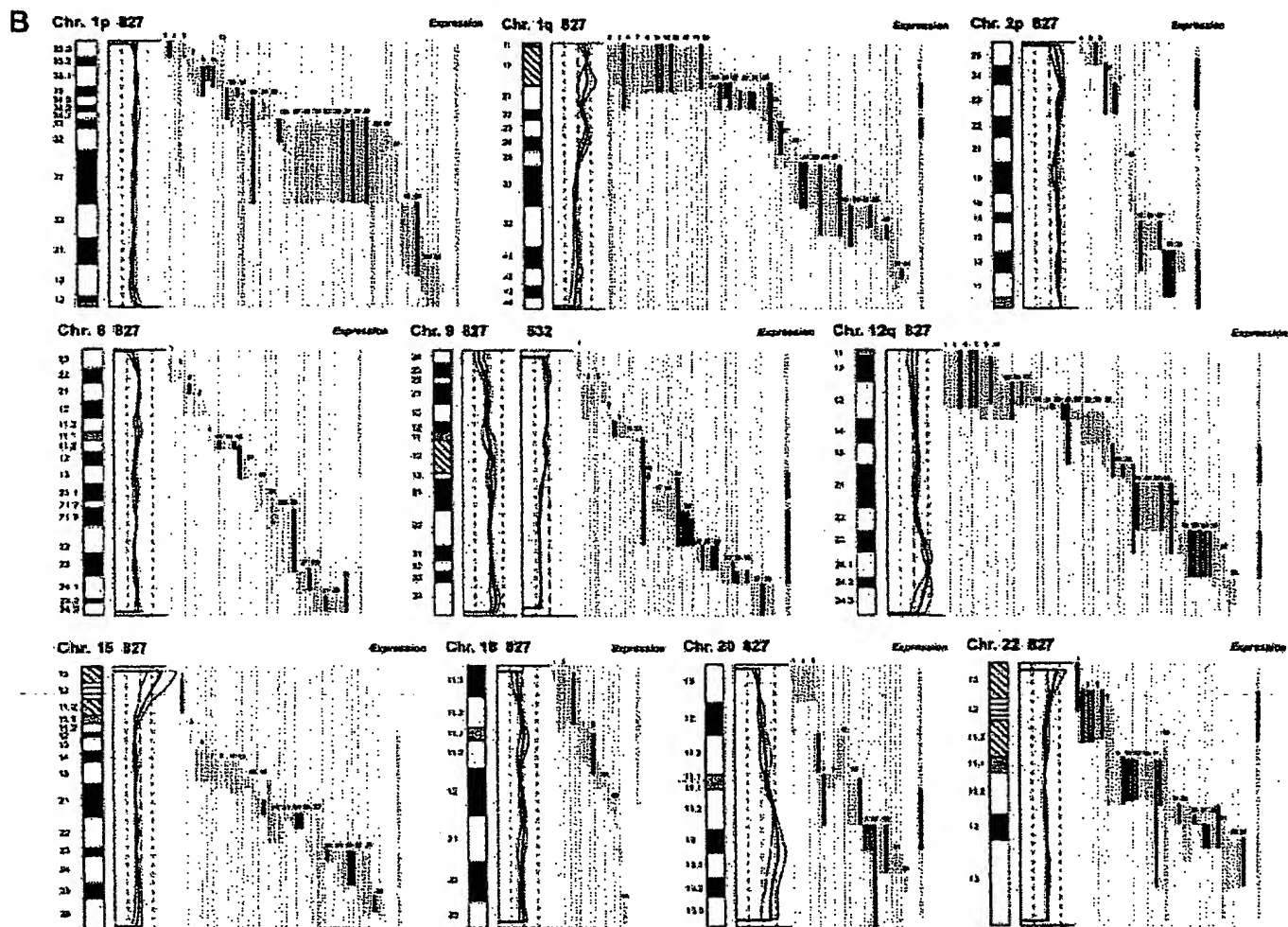


FIG. 1—continued

for 30 min at 25 °C followed by 10 washes in 6× SSPE-T at 25 °C. The probe arrays were scanned at 560 nm using a confocal laser scanning microscope (made for Affymetrix by Hewlett-Packard). The readings from the quantitative scanning were analyzed by Affymetrix gene expression analysis software.

Microsatellite Analysis—Microsatellite Analysis was performed as described previously (14). Microsatellites were selected by use of www.ncbi.nlm.nih.gov/genemap98, and primer sequences were obtained from the genome data base at www.gdb.org. DNA was extracted from tumor and blood and amplified by PCR in a volume of 20 μ l for 35 cycles. The amplicons were denatured and electrophoresed for 3 h in an ABI Prism 377. Data were collected in the Gene Scan program for fragment analysis. Loss of heterozygosity was defined as less than 33% of one allele detected in tumor amplicons compared with blood.

Proteomic Analysis—TCCs were minced into small pieces and homogenized in a small glass homogenizer in 0.5 ml of lysis solution. Samples were stored at –20 °C until use. The procedure for 2D gel electrophoresis has been described in detail elsewhere (15, 16). Gels were stained with silver nitrate and/or Coomassie Brilliant Blue. Proteins were identified by a combination of procedures that included microsequencing, mass spectrometry, two-dimensional gel Western immunoblotting, and comparison with the master two-dimensional gel image of human keratinocyte proteins; see biobase.dk/cgi-bin/celis.

CGH—Hybridization of differentially labeled tumor and normal DNA to normal metaphase chromosomes was performed as described previously (10). Fluorescein-labeled tumor DNA (200 ng), Texas Red-

labeled reference DNA (200 ng), and human Cot-1 DNA (20 μ g) were denatured at 37 °C for 5 min and applied to denatured normal metaphase slides. Hybridization was at 37 °C for 2 days. After washing, the slides were counterstained with 0.15 μ g/ml 4,6-diamidino-2-phenylindole in an anti-fade solution. A second hybridization was performed for all tumor samples using fluorescein-labeled reference DNA and Texas Red-labeled tumor DNA (inverse labeling) to confirm the aberrations detected during the initial hybridization. Each CGH experiment also included a normal control hybridization using fluorescein- and Texas Red-labeled normal DNA. Digital image analysis was used to identify chromosomal regions with abnormal fluorescence ratios, indicating regions of DNA gains and losses. The average green:red fluorescence intensity ratio profiles were calculated using four images of each chromosome (eight chromosomes total) with normalization of the green:red fluorescence intensity ratio for the entire metaphase and background correction. Chromosome identification was performed based on 4,6-diamidino-2-phenylindole banding patterns. Only images showing uniform high intensity fluorescence with minimal background staining were analyzed. All centromeres, p arms of acrocentric chromosomes, and heterochromatic regions were excluded from the analysis.

RESULTS

Comparative Genomic Hybridization—The CGH analysis identified a number of chromosomal gains and losses in the

Gene Copy Numbers, Transcripts, and Protein Levels

TABLE I
Correlation between alterations detected by CGH and by expression monitoring

Top, CGH used as independent variable (if CGH alteration – what expression ratio was found); bottom, altered expression used as independent variable (if expression alteration – what CGH deviation was found).

CGH alterations	Tumor 733 vs. 335	Concordance	CGH alterations	Tumor 827 vs. 532	Concordance
	Expression change clusters			Expression change clusters	
13 Gain	10 Up-regulation 0 Down-regulation 3 No change	77%	10 Gain	8 Up-regulation 0 Down-regulation 2 No change	80%
10 Loss	1 Up-regulation 5 Down-regulation 4 No change	50%	12 Loss	3 Up-regulation 2 Down regulation 7 No change	17%
Expression change clusters	Tumor 733 vs. 335	Concordance	Expression change clusters	Tumor 827 vs. 532	Concordance
	CGH alterations			CGH alterations	
16 Up-regulation	11 Gain 2 Loss 3 No change	69%	17 Up-regulation	10 Gain 5 Loss 2 No change	59%
21 Down-regulation	1 Gain 8 Loss 12 No change	38%	9 Down-regulation	0 Gain 3 Loss 6 No change	33%
15 No change	3 Gain 3 Loss 9 No change	60%	21 No change	1 Gain 3 Loss 17 No change	81%

two invasive tumors (stage pT1, TCCs 733 and 827), whereas the two non-invasive papillomas (stage pTa, TCCs 335 and 532) showed only 9p–, 9q22–q33–, and X–, and 7+, 9q–, and Y–, respectively. Both invasive tumors showed changes (1q22–24+, 2q14.1–qter–, 3q12–q13.3–, 6q12–q22–, 9q34+, 11q12–q13+, 17+, and 20q11.2–q12+) that are typical for their disease stage, as well as additional alterations, some of which are shown in Fig. 1. Areas with gains and losses deviated from the normal copy number to some extent, and the average numerical deviation from normal was 0.4-fold in the case of TCC 733 and 0.3-fold for TCC 827. The largest changes, amounting to at least a doubling of chromosomal content, were observed at 1q23 in TCC 733 (Fig. 1A) and 20q12 in TCC 827 (Fig. 1B).

mRNA Expression in Relation to DNA Copy Number—The mRNA levels from the two invasive tumors (TCCs 827 and 733) were compared with the two non-invasive counterparts (TCCs 532 and 335). This was done in two separate experiments in which we compared TCCs 733 to 335 and 827 to 532, respectively, using two different scaling settings for the arrays to rule out scaling as a confounding parameter. Approximately 1,800 genes that yielded a signal on the arrays were searched in the Unigene and Genemap data bases for chromosomal location, and those with a known location (1096) were plotted as bars covering their purported locus. In that way it was possible to construct a graphic presentation of DNA copy number and relative mRNA levels along the individual chromosomes (Fig. 1).

For each mRNA a ratio was calculated between the level in the invasive versus the non-invasive counterpart. Bars, which represent chromosomal location of a gene, were color-coded according to the expression ratio, and only differences larger

than 2-fold were regarded as informative (Fig. 1). The density of genes along the chromosomes varied, and areas containing only one gene were excluded from the calculations. The resolution of the CGH method is very low, and some of the outlier data may be because of the fact that the boundaries of the chromosomal aberrations are not known at high resolution.

Two sets of calculations were made from the data. For the first set we used CGH alterations as the independent variable and estimated the frequency of expression alterations in these chromosomal areas. In general, areas with a strong gain of chromosomal material contained a cluster of genes having increased mRNA expression. For example, both chromosomes 1q21–q25, 2p and 9q, showed a relative gain of more than 100% in DNA copy number that was accompanied by increased mRNA expression levels in the two tumor pairs (Fig. 1). In most cases, chromosomal gains detected by CGH were accompanied by an increased level of transcripts in both TCCs 733 (77%) and 827 (80%) (Table I, top). Chromosomal losses, on the other hand, were not accompanied by decreased expression in several cases, and were often registered as having unaltered RNA levels (Table I, top). The inability to detect RNA expression changes in these cases was not because of fewer genes mapping to the lost regions (data not shown).

In the second set of calculations we selected expression alterations above 2-fold as the independent variable and estimated the frequency of CGH alterations in these areas. As above, we found that increased transcript expression correlated with gain of chromosomal material (TCC 733, 69% and TCC 827, 59%), whereas reduced expression was often detected in areas with unaltered CGH ratios (Table I, bottom). Furthermore, as a control we looked at areas with no alter-

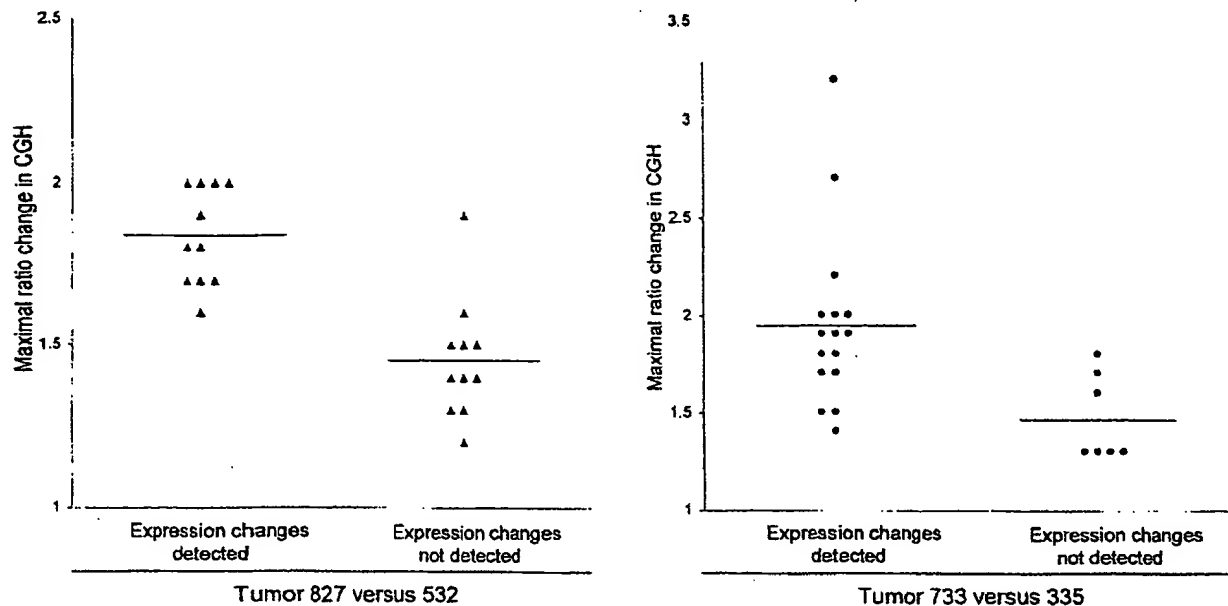


FIG. 2. Correlation between maximum CGH aberration and the ability to detect expression change by oligonucleotide array monitoring. The aberration is shown as a numerical -fold change in ratio between invasive tumors 827 (▲) and 733 (◆) and their non-invasive counterparts 532 and 335. The expression change was taken from the *Expression* line to the right in Fig. 1, which depicts the resulting expression change for a given chromosomal region. At least half of the mRNAs from a given region have to be either up- or down-regulated to be scored as an expression change. All chromosomal arms in which the CGH ratio plus or minus one standard deviation was outside the ratio value of one were included.

ation in expression. No alteration was detected by CGH in most of these areas (TCC 733, 60% and TCC 827, 81%; see Table I, *bottom*). Because the ability to observe reduced or increased mRNA expression clustering to a certain chromosomal area clearly reflected the extent of copy number changes, we plotted the maximum CGH aberrations in the regions showing CGH changes against the ability to detect a change in mRNA expression as monitored by the oligonucleotide arrays (Fig. 2). For both tumors TCC 733 ($p < 0.015$) and TCC 827 ($p < 0.00003$) a highly significant correlation was observed between the level of CGH ratio change (reflecting the DNA copy number) and alterations detected by the array based technology (Fig. 2). Similar data were obtained when areas with altered expression were used as independent variables. These areas correlated best with CGH when the CGH ratio deviated 1.6- to 2.0-fold (Table I, *bottom*) but mostly did not at lower CGH deviations. These data probably reflect that loss of an allele may only lead to a 50% reduction in expression level, which is at the cut-off point for detection of expression alterations. Gain of chromosomal material can occur to a much larger extent.

Microsatellite-based Detection of Minor Areas of Losses—In TCC 733, several chromosomal areas exhibiting DNA amplification were preceded or followed by areas with a normal CGH but reduced mRNA expression (see Fig. 1, TCC 733 chromosome 1q32, 2p21, and 7q21 and q32, 9q34, and 10q22). To determine whether these results were because of undetected loss of chromosomal material in these regions or

because of other non-structural mechanisms regulating transcription, we examined two microsatellites positioned at chromosome 1q25–32 and two at chromosome 2p22. Loss of heterozygosity (LOH) was found at both 1q25 and at 2p22 indicating that minor deleted areas were not detected with the resolution of CGH (Fig. 3). Additionally, chromosome 2p in TCC 733 showed a CGH pattern of gain/no change/gain of DNA that correlated with transcript increase/decrease/increase. Thus, for the areas showing increased expression there was a correlation with the DNA copy number alterations (Fig. 1A). As indicated above, the mRNA decrease observed in the middle of the chromosomal gain was because of LOH, implying that one of the mechanisms for mRNA down-regulation may be regions that have undergone smaller losses of chromosomal material. However, this cannot be detected with the resolution of the CGH method.

In both TCC 733 and TCC 827, the telomeric end of chromosome 11p showed a normal ratio in the CGH analysis; however, clusters of five and three genes, respectively, lost their expression. Two microsatellites (D11S1760, D11S922) positioned close to MUC2, IGF2, and cathepsin D indicated LOH as the most likely mechanism behind the loss of expression (data not shown).

A reduced expression of mRNA observed in TCC 733 at chromosomes 3q24, 11p11, 12p12.2, 12q21.1, and 16q24 and in TCC 827 at chromosome 11p15.5, 12p11, 15q11.2, and 18q12 was also examined for chromosomal losses using microsatellites positioned as close as possible to the gene loci

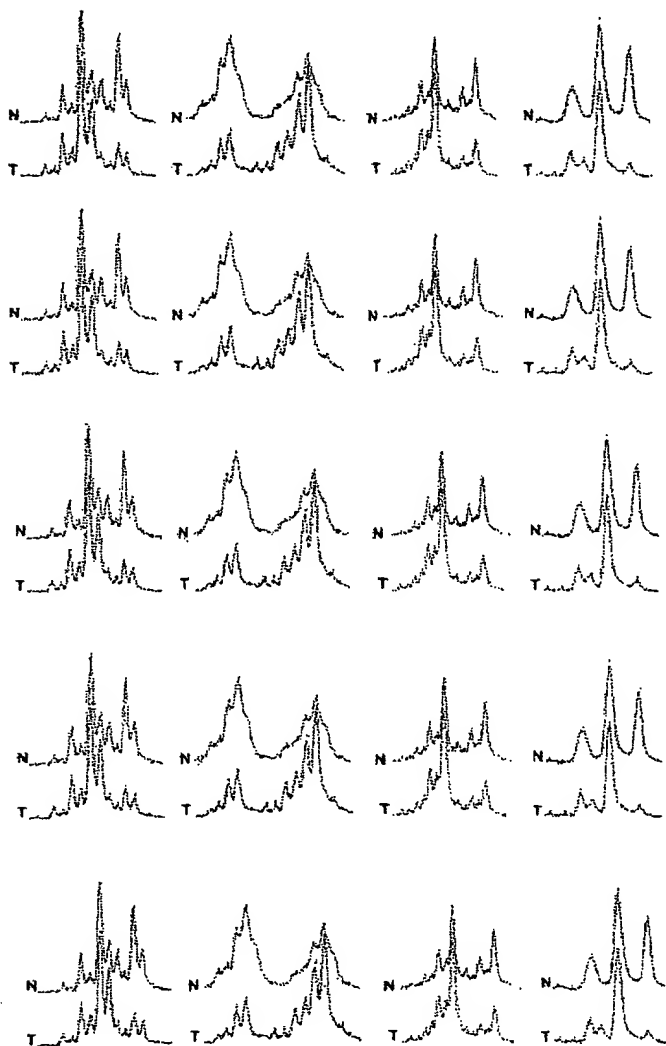


FIG. 3. Microsatellite analysis of loss of heterozygosity. Tumor 733 showing loss of heterozygosity at chromosome 1q25, detected (a) by D1S215 close to Hu class I histocompatibility antigen (gene number 38 in Fig. 1), (b) by D1S2735 close to cathepsin E (gene number 41 in Fig. 1), and (c) at chromosome 2p23 by D2S2251 close to general β -spectrin (gene number 11 on Fig. 1) and of (d) tumor 827 showing loss of heterozygosity at chromosome 18q12 by S18S1118 close to mitochondrial 3-oxoacyl-coenzyme A thiolase (gene number 12 in Fig. 1). The upper curves show the electropherogram obtained from normal DNA from leukocytes (N), and the lower curves show the electropherogram from tumor DNA (T). In all cases one allele is partially lost in the tumor amplicon.

showing reduced mRNA transcripts. Only the microsatellite positioned at 18q12 showed LOH (Fig. 3), suggesting that transcriptional down-regulation of genes in the other regions may be controlled by other mechanisms.

Relation between Changes in mRNA and Protein Levels—2D-PAGE analysis, in combination with Coomassie Brilliant Blue and/or silver staining, was carried out on all four tumors using fresh biopsy material. 40 well resolved abundant known proteins migrating in areas away from the edges of the pH

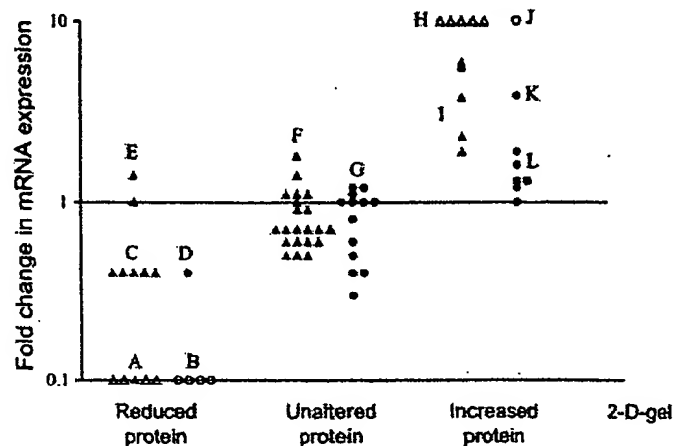


FIG. 4. Correlation between protein levels as judged by 2D-PAGE and transcript ratio. For comparison proteins were divided in three groups, unaltered in level or up- or down-regulated (horizontal axis). The mRNA ratio as determined by oligonucleotide arrays was plotted for each gene (vertical axis). ▲, mRNAs that were scored as present in both tumors used for the ratio calculation; △, mRNAs that were scored as absent in the invasive tumors (along horizontal axis) or as absent in non-invasive reference (top of figure). Two different scalings were used to exclude scaling as a confounder, TCCs 827 and 532 (▲△) were scaled with background suppression, and TCCs 733 and 335 (●○) were scaled without suppression. Both comparisons showed highly significant ($p < 0.005$) differences in mRNA ratios between the groups. Proteins shown, were as follows: Group A (from left), phosphoglucomutase 1, glutathione transferase class μ number 4, fatty acid-binding protein homologue, cytochrome 15, and cytochrome 13; B (from left), fatty acid-binding protein homologue, 28-kDa heat shock protein, cytochrome 13, and calnexin; C (from left), α -enolase, hnRNP B1, 28-kDa heat shock protein, 14-3-3- ϵ , and pre-mRNA splicing factor; D, mesothelial keratin K7 (type II); E (from top), glutathione S-transferase- π and mesothelial keratin K7 (type II); F (from top and left), adenyl cyclase-associated protein, E-cadherin, keratin 19, calgizzarin, phosphoglycerate mutase, annexin IV, cytoskeletal γ -actin, hnRNP A1, integral membrane protein calnexin (IP90), hnRNP H, brain-type clathrin light chain- α , hnRNP F, 70-kDa heat shock protein, heterogeneous nuclear ribonucleoprotein A/B, translationally controlled tumor protein, liver glyceraldehyde-3-phosphate dehydrogenase, keratin 8, aldehyde reductase, and Na,K-ATPase β -1 subunit; G, (from top and left), TCP20, calgizzarin, 70-kDa heat shock protein, calnexin, hnRNP H, cytochrome 15, ATP synthase, keratin 19, triosephosphate isomerase, hnRNP F, liver glyceraldehyde-3-phosphatase dehydrogenase, glutathione S-transferase- π , and keratin 8; H (from left), plasma gelsolin, autoantigen calreticulin, thioredoxin, and NAD $^{+}$ -dependent 15 hydroxyprostaglandin dehydrogenase; I (from top), prollyl 4-hydroxylase β -subunit, cytochrome 20, cytochrome 17, prohibition, and fructose 1,6-bisphosphatase; J annexin II; K, annexin IV; L (from top and left), 90-kDa heat shock protein, prollyl 4-hydroxylase β -subunit, α -enolase, GRP 78, cyclophilin, and cofilin.

gradient, and having a known chromosomal location, were selected for analysis in the TCC pair 827/532. Proteins were identified by a combination of methods (see "Experimental Procedures"). In general there was a highly significant correlation ($p < 0.005$) between mRNA and protein alterations (Fig. 4). Only one gene showed disagreement between transcript alteration and protein alteration. Except for a group of cyto-

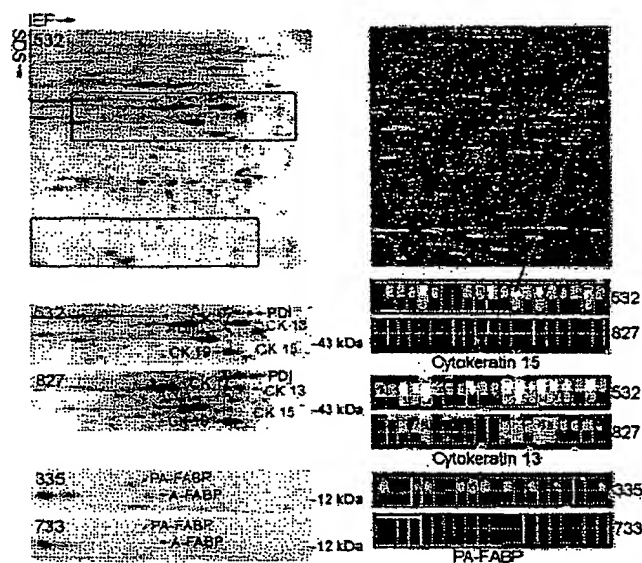


FIG. 5. Comparison of protein and transcript levels in invasive and non-invasive TCCs. The upper part of the figure shows a 2D gel (left) and the oligonucleotide array (right) of TCC 532. The red rectangles on the upper gel highlight the areas that are compared below. Identical areas of 2D gels of TCCs 532 and 827 are shown below. Clearly, cytokeratins 13 and 15 are strongly down-regulated in TCC 827 (red annotation). The tile on the array containing probes for cytokeratin 15 is enlarged below the array (red arrow) from TCC 532 and is compared with TCC 827. The upper row of squares in each tile corresponds to perfect match probes; the lower row corresponds to mismatch probes containing a mutation (used for correction for unspecific binding). Absence of signal is depicted as black, and the higher the signal the lighter the color. A high transcript level was detected in TCC 532 (6151 units) whereas a much lower level was detected in TCC 827 (absence of signals). For cytokeratin 13, a high transcript level was also present in TCC 532 (15659 units), and a much lower level was present in TCC 827 (623 units). The 2D gels at the bottom of the figure (left) show levels of PA-FABP and adipocyte-FABP in TCCs 335 and 733 (invasive), respectively. Both proteins are down-regulated in the invasive tumor. To the right we show the array tiles for the PA-FABP transcript. A medium transcript level was detected in the case of TCC 335 (1277 units) whereas very low levels were detected in TCC 733 (166 units). IEF, isoelectric focusing.

keratins encoded by genes on chromosome 17 (Fig. 5) the analyzed proteins did not belong to a particular family. 26 well focused proteins whose genes had a known chromosomal location were detected in TCCs 733 and 335, and of these 19 correlated ($p < 0.005$) with the mRNA changes detected using the arrays (Fig. 4). For example, PA-FABP was highly expressed in the non-invasive TCC 335 but lost in the invasive counterpart (TCC 733; see Fig. 5). The smaller number of proteins detected in both 733 and 335 was because of the smaller size of the biopsies that were available.

11 chromosomal regions where CGH showed aberrations that corresponded to the changes in transcript levels also showed corresponding changes in the protein level (Table II). These regions included genes that encode proteins that are found to be frequently altered in bladder cancer, namely cytokeratins 17 and 20, annexins II and IV, and the fatty acid-binding proteins PA-FABP and FABP1. Four of these proteins were encoded by genes in chromosome 17q, a frequently amplified chromosomal area in invasive bladder cancers.

DISCUSSION

Most human cancers have abnormal DNA content, having lost some chromosomal parts and gained others. The present study provides some evidence as to the effect of these gains and losses on gene expression in two pairs of non-invasive and invasive TCCs using high throughput expression arrays and proteomics, in combination with CGH. In general, the results showed that there is a clear individual regulation of the mRNA expression of single genes, which in some cases was superimposed by a DNA copy number effect. In most cases, genes located in chromosomal areas with gains often exhibited increased mRNA expression, whereas areas showing losses showed either no change or a reduced mRNA expression. The latter might be because of the fact that losses most often are restricted to loss of one allele, and the cut-off point for detection of expression alterations was a 2-fold change, thus being at the border of detection. In several cases, how-

TABLE II
Proteins whose expression level correlates with both mRNA and gene dose changes

Protein	Chromosomal location	Tumor TCC	CGH alteration	Transcript alteration ^a	Protein alteration
Annexin II	1q21	733	Gain	Abs to Pres ^a	Increase
Annexin IV	2p13	733	Gain	3.9-Fold up	Increase
Cytokeratin 17	17q12-q21	827	Gain	3.8-Fold up	Increase
Cytokeratin 20	17q21.1	827	Gain	5.6-Fold up	Increase
(PA)-FABP	8q21.2	827	Loss	10-Fold down	Decrease
FABP1	9q22	827	Gain	2.3-Fold up	Increase
Plasma gelsolin	9q31	827	Gain	Abs to Pres	Increase
Heat shock protein 28	15q12-q13	827	Loss	2.5-Fold up	Decrease
Prohibitin	17q21	827/733	Gain	3.7-/2.5-Fold up ^b	Increase
Prolyl-4-hydroxyl	17q25	827/733	Gain	5.7-/1.6-Fold up	Increase
hnRNPB1	7p15	827	Loss	2.5-Fold down	Decrease

^a Abs, absent; Pres, present.

^b In cases where the corresponding alterations were found in both TCCs 827 and 733 these are shown as 827/733.

ever, an increase or decrease in DNA copy number was associated with *de novo* occurrence or complete loss of transcript, respectively. Some of these transcripts could not be detected in the non-invasive tumor but were present at relatively high levels in areas with DNA amplifications in the invasive tumors (e.g. in TCC 733 transcript from cellular ligand of annexin II gene (chromosome 1q21) from absent to 2670 arbitrary units; in TCC 827 transcript from small proline-rich protein 1 gene (chromosome 1q12-q21.1) from absent to 1326 arbitrary units). It may be anticipated from these data that significant clustering of genes with an increased expression to a certain chromosomal area indicates an increased likelihood of gain of chromosomal material in this area.

Considering the many possible regulatory mechanisms acting at the level of transcription, it seems striking that the gene dose effects were so clearly detectable in gained areas. One hypothetical explanation may lie in the loss of controlled methylation in tumor cells (17–19). Thus, it may be possible that in chromosomes with increased DNA copy numbers two or more alleles could be demethylated simultaneously leading to a higher transcription level, whereas in chromosomes with losses the remaining allele could be partly methylated, turning off the process (20, 21). A recent report has documented a ploidy regulation of gene expression in yeast, but in this case all the genes were present in the same ratio (22), a situation that is not analogous to that of cancer cells, which show marked chromosomal aberrations, as well as gene dosage effects.

Several CGH studies of bladder cancer have shown that some chromosomal aberrations are common at certain stages of disease progression, often occurring in more than 1 of 3 tumors. In pTa tumors, these include 9p–, 9q–, 1q+, Y– (2, 6), and in pT1 tumors, 2q–, 11p–, 11q–, 1q+, 5p+, 8q+, 17q+, and 20q+ (2–4, 6, 7). The pTa tumors studied here showed similar aberrations such as 9p– and 9q22–q33– and 9q– and Y–, respectively. Likewise, the two minimal invasive pT1 tumors showed aberrations that are commonly seen at that stage, and TCC 827 had a remarkable resemblance to the commonly seen pattern of losses and gains, such as 1q22–24 amplification (seen in both tumors), 11q14–q22 loss, the latter often linked to 17 q+ (both tumors), and 1q+ and 9p–, often linked to 20q+ and 11 q13+ (both tumors) (7–9). These observations indicate that the pairs of tumors used in this study exhibit chromosomal changes observed in many tumors, and therefore the findings could be of general importance for bladder cancer.

Considering that the mapping resolution of CGH is of about 20 megabases it is only possible to get a crude picture of chromosomal instability using this technique. Occasionally, we observed reduced transcript levels close to or inside regions with increased copy numbers. Analysis of these regions by positioning heterozygous microsatellites as close as possible to the locus showing reduced gene expression revealed loss of heterozygosity in several cases. It seems likely that multiple and different events occur along each chromosomal

arm and that the use of cDNA microarrays for analysis of DNA copy number changes will reach a resolution that can resolve these changes, as has recently been proposed (2). The outlier data were not more frequent at the boundaries of the CGH aberrations. At present we do not know the mechanism behind chromosomal aneuploidy and cannot predict whether chromosomal gains will be transcribed to a larger extent than the two native alleles. A mechanism as genetic imprinting has an impact on the expression level in normal cells and is often reduced in tumors. However, the relation between imprinting and gain of chromosomal material is not known.

We regard it as a strength of this investigation that we were able to compare invasive tumors to benign tumors rather than to normal urothelium, as the tumors studied were biologically very close and probably may represent successive steps in the progression of bladder cancer. Despite the limited amount of fresh tissue available it was possible to apply three different state of the art methods. The observed correlation between DNA copy number and mRNA expression is remarkable when one considers that different pieces of the tumor biopsies were used for the different sets of experiments. This indicates that bladder tumors are relatively homogenous, a notion recently supported by CGH and LOH data that showed a remarkable similarity even between tumors and distant metastasis (10, 23).

In the few cases analyzed, mRNA and protein levels showed a striking correspondence although in some cases we found discrepancies that may be attributed to translational regulation, post-translational processing, protein degradation, or a combination of these. Some transcripts belong to undertranslated mRNA pools, which are associated with few translationally inactive ribosomes; these pools, however, seem to be rare (24). Protein degradation, for example, may be very important in the case of polypeptides with a short half-life (e.g. signaling proteins). A poor correlation between mRNA and protein levels was found in liver cells as determined by arrays and 2D-PAGE (25), and a moderate correlation was recently reported by Ideker *et al.* (26) in yeast.

Interestingly, our study revealed a much better correlation between gained chromosomal areas and increased mRNA levels than between loss of chromosomal areas and reduced mRNA levels. In general, the level of CGH change determined the ability to detect a change in transcript. One possible explanation could be that by losing one allele the change in mRNA level is not so dramatic as compared with gain of material, which can be rather unlimited and may lead to a severalfold increase in gene copy number resulting in a much higher impact on transcript level. The latter would be much easier to detect on the expression arrays as the cut-off point was placed at a 2-fold level so as not to be biased by noise on the array. Construction of arrays with a better signal to noise ratio may in the future allow detection of lesser than 2-fold alterations in transcript levels, a feature that may facilitate the analysis of the effect of loss of chromosomal areas on transcript levels.

In eleven cases we found a significant correlation between DNA copy number, mRNA expression, and protein level. Four of these proteins were encoded by genes located at a frequently amplified area in chromosome 17q. Whether DNA copy number is one of the mechanisms behind alteration of these eleven proteins is at present unknown and will have to be proved by other methods using a larger number of samples. One factor making such studies complicated is the large extent of protein modification that occurs after translation, requiring immunoidentification and/or mass spectrometry to correctly identify the proteins in the gels.

In conclusion, the results presented in this study exemplify the large body of knowledge that may be possible to gather in the future by combining state of the art techniques that follow the pathway from DNA to protein (26). Here, we used a traditional chromosomal CGH method, but in the future high resolution CGH based on microarrays with many thousand radiation hybrid-mapped genes will increase the resolution and information derived from these types of experiments (2). Combined with expression arrays analyzing transcripts derived from genes with known locations, and 2D gel analysis to obtain information at the post-translational level, a clearer and more developed understanding of the tumor genome will be forthcoming.

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TECHNICAL UPDATE

FROM YOUR LABORATORY SERVICES PROVIDER

HER-2/neu Breast Cancer Predictive Testing

Julie Sanford Hanna, Ph.D. and Dan Mornin, M.D.

EACH YEAR, OVER 182,000 WOMEN in the United States are diagnosed with breast cancer, and approximately 45,000 die of the disease.¹ Incidence appears to be increasing in the United States at a rate of roughly 2% per year. The reasons for the increase are unclear, but non-genetic risk factors appear to play a large role.²

Five-year survival rates range from approximately 65%-85%, depending on demographic group, with a significant percentage of women experiencing recurrence of their cancer within 10 years of diagnosis. One of the factors most predictive for recurrence once a diagnosis of breast cancer has been made is the number of axillary lymph nodes to which tumor has metastasized. Most node-positive women are given adjuvant therapy, which increases their survival. However, 20%-30% of patients without axillary node involvement also develop recurrent disease, and the difficulty lies in how to identify this high-risk subset of patients. These patients could benefit from increased surveillance, early intervention, and treatment.

Prognostic markers currently used in breast cancer recurrence prediction include tumor size, histological grade, steroid hormone receptor status, DNA ploidy, proliferative index, and cathepsin D status. Expression of growth factor receptors and over-expression of the HER-2/neu oncogene have also been identified as having value regarding treatment regimen and prognosis.

HER-2/neu (also known as c-erbB2) is an oncogene that encodes a transmembrane glycoprotein that is homologous to, but distinct from, the epidermal growth factor receptor. Numerous studies have indicated that high levels of expression of this protein are associated with rapid tumor growth, certain forms of therapy resistance, and shorter disease-free survival. The gene has been shown to be amplified and/or overexpressed in 10%-30% of invasive breast cancers and in 40%-60% of intraductal breast carcinoma.³

There are two distinct FDA-approved methods by which HER-2/neu status can be evaluated: immunohistochemistry (IHC, HercepTest™) and FISH (fluorescent in situ hybridization, PathVysion™ Kit). Both methods can be performed on archived and current specimens. The first method allows visual assessment of the amount of HER-2/neu protein present on the cell membrane. The latter method allows direct quantification of the level of gene amplification present in the tumor, enabling differentiation between low- versus high-amplification. At least one study has demonstrated a difference in

recurrence risk in women younger than 40 years of age for low- versus high-amplified tumors (54.5% compared to 85.7%); this is compared to a recurrence rate of 16.7% for patients with no HER-2/neu gene amplification.⁴ HER-2/neu status may be particularly important to establish in women with small (≤ 1 cm) tumor size.

The choice of methodology for determination of HER-2/neu status depends in part on the clinical setting. FDA approval for the Vysis FISH test was granted based on clinical trials involving 1549 node-positive patients. Patients received one of three different treatments consisting of different doses of cyclophosphamide, Adriamycin, and 5-fluorouracil (CAF). The study showed that patients with amplified HER-2/neu benefited from treatment with higher doses of adriamycin-based therapy, while those with normal HER-2/neu levels did not. The study therefore identified a sub-set of women, who because they did not benefit from more aggressive treatment, did not need to be exposed to the associated side effects. In addition, other evidence indicates that HER-2/neu amplification in node-negative patients can be used as an independent prognostic indicator for early recurrence, recurrent disease at any time and disease-related death.⁵ Demonstration of HER-2/neu gene amplification by FISH has also been shown to be of value in predicting response to chemotherapy in stage-2 breast cancer patients.

Selection of patients for Herceptin® (Trastuzumab) monoclonal antibody therapy, however, is based upon demonstration of HER-2/neu protein overexpression using HercepTest™. Studies using Herceptin® in patients with metastatic breast cancer show an increase in time to disease progression, increased response rate to chemotherapeutic agents and a small increase in overall survival rate. The FISH assays have not yet been approved for this purpose, and studies looking at response to Herceptin® in patients with or without gene amplification status determined by FISH are in progress.

In general, FISH and IHC results correlate well. However, subsets of tumors are found which show discordant results; i.e., protein overexpression without gene amplification or lack of protein overexpression with gene amplification. The clinical significance of such results is unclear. Based on the above considerations, HER-2/neu testing at SHMC/PAML will utilize immunohistochemistry (HercepTest®) as a screen, followed by FISH in IHC-negative cases. Alternatively, either method may be ordered individually depending on the clinical setting or clinician preference.

CPT code information

HER-2/neu via IHC

88342 (including interpretive report)

HER-2/neu via FISH

- 8827 1x2 Molecular cytogenetics, DNA probe, each
88274 Molecular cytogenetics, interphase in situ hybridization, analyze 25-99 cells
88291 Cytogenetics and molecular cytogenetics, interpretation and report

Procedural Information

Immunohistochemistry is performed using the FDA-approved DAKO antibody kit, Herceptest[®]. The DAKO kit contains reagents required to complete a two-step immunohistochemical staining procedure for routinely processed, paraffin-embedded specimens. Following incubation with the primary rabbit antibody to human HER-2/neu protein, the kit employs a ready-to-use dextran-based visualization reagent. This reagent consists of both secondary goat anti-rabbit antibody molecules with horseradish peroxidase molecules linked to a common dextran polymer backbone, thus eliminating the need for sequential application of link antibody and peroxidase conjugated antibody. Enzymatic conversion of the subsequently added chromogen results in formation of visible reaction product at the antigen site. The specimen is then counterstained; a pathologist using light-microscopy interprets results.

FISH analysis at SHMC/PAML is performed using the FDA-approved PathVysion[™] HER-2/neu DNA probe kit, produced by Vysis, Inc. Formalin fixed, paraffin-embedded breast tissue is processed using routine histological methods, and then slides are treated to allow hybridization of DNA probes to the nuclei present in the tissue section. The Pathvysion[™] kit contains two direct-labeled DNA probes, one specific for the alphoid repetitive DNA (CEP 17, spectrum orange) present at the chromosome 17 centromere and the second for the HER-2/neu oncogene located at 17q11.2-12 (spectrum green). Enumeration of the probes allows a ratio of the number of copies of chromosome 17 to the number of copies of HER-2/neu to be obtained; this enables quantification of low versus high amplification levels, and allows an estimate of the percentage of cells with HER-2/neu gene amplification. The clinically relevant distinction is whether the gene amplification is due to increased gene copy number on the two chromosome 17 homologues normally present or an increase in the number of chromosome 17s in the cells. In the majority of cases, ratio equivalents less than 2.0 are indicative of a normal/negative result, ratios of 2.1 and over indicate that amplification is present and to what degree. Interpretation of this data will be performed and reported from the Vysis-certified Cytogenetics laboratory at SHMC.

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SECOND DECLARATION OF PAUL POLAKIS, Ph.D.

I, Paul Polakis, Ph.D., declare and say as follows:

1. I am currently employed by Genentech, Inc. where my job title is Staff Scientist.
2. Since joining Genentech in 1999, one of my primary responsibilities has been leading Genentech's Tumor Antigen Project, which is a large research project with a primary focus on identifying tumor cell markers that find use as targets for both the diagnosis and treatment of cancer in humans.
3. As I stated in my previous Declaration dated May 7, 2004 (attached as Exhibit A), my laboratory has been employing a variety of techniques, including microarray analysis, to identify genes which are differentially expressed in human tumor tissue relative to normal human tissue. The primary purpose of this research is to identify proteins that are abundantly expressed on certain human tumor tissue(s) and that are either (i) not expressed, or (ii) expressed at detectably lower levels, on normal tissue(s).
4. In the course of our research using microarray analysis, we have identified approximately 200 gene transcripts that are present in human tumor tissue at significantly higher levels than in normal human tissue. To date, we have successfully generated antibodies that bind to 31 of the tumor antigen proteins expressed from these differentially expressed gene transcripts and have used these antibodies to quantitatively determine the level of production of these tumor antigen proteins in both human tumor tissue and normal tissue. We have then quantitatively compared the levels of mRNA and protein in both the tumor and normal tissues analyzed. The results of these analyses are attached herewith as Exhibit B. In Exhibit B, "+" means that the mRNA or protein was detectably overexpressed in the tumor tissue relative to normal tissue and "-" means that no detectable overexpression was observed in the tumor tissue relative to normal tissue.
5. As shown in Exhibit B, of the 31 genes identified as being detectably overexpressed in human tumor tissue as compared to normal human tissue at the mRNA level, 28 of them (i.e., greater than 90%) are also detectably overexpressed in human tumor tissue as compared to normal human tissue at the protein level. As such, in the cases where we have been able to quantitatively measure both (i) mRNA and (ii) protein levels in both (i) tumor tissue and (ii) normal tissue, we have observed that in the vast majority of cases, there is a very strong correlation between increases in mRNA expression and increases in the level of protein encoded by that mRNA.

6. Based upon my own experience accumulated in more than 20 years of research, including the data discussed in paragraphs 4-5 above and my knowledge of the relevant scientific literature, it is my considered scientific opinion that for human genes, an increased level of mRNA in a tumor tissue relative to a normal tissue more often than not correlates to a similar increase in abundance of the encoded protein in the tumor tissue relative to the normal tissue. In fact, it remains a generally accepted working assumption in molecular biology that increased mRNA levels are more often than not predictive of elevated levels of the encoded protein. In fact, an entire industry focusing on the research and development of therapeutic antibodies to treat a variety of human diseases, such as cancer, operates on this working assumption.
7. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: 3-29-00

By: Paul Polakis

Paul Polakis, Ph.D.

EXHIBIT A

DECLARATION OF PAUL POLAKIS, Ph.D.

I, Paul Polakis, Ph.D., declare and say as follows:

1. I was awarded a Ph.D. by the Department of Biochemistry of the Michigan State University in 1984. My scientific Curriculum Vitae is attached to and forms part of this Declaration (Exhibit A).
2. I am currently employed by Genentech, Inc. where my job title is Staff Scientist. Since joining Genentech in 1999, one of my primary responsibilities has been leading Genentech's Tumor Antigen Project, which is a large research project with a primary focus on identifying tumor cell markers that find use as targets for both the diagnosis and treatment of cancer in humans.
3. As part of the Tumor Antigen Project, my laboratory has been analyzing differential expression of various genes in tumor cells relative to normal cells. The purpose of this research is to identify proteins that are abundantly expressed on certain tumor cells and that are either (i) not expressed, or (ii) expressed at lower levels, on corresponding normal cells. We call such differentially expressed proteins "tumor antigen proteins". When such a tumor antigen protein is identified, one can produce an antibody that recognizes and binds to that protein. Such an antibody finds use in the diagnosis of human cancer and may ultimately serve as an effective therapeutic in the treatment of human cancer.
4. In the course of the research conducted by Genentech's Tumor Antigen Project, we have employed a variety of scientific techniques for detecting and studying differential gene expression in human tumor cells relative to normal cells, at genomic DNA, mRNA and protein levels. An important example of one such technique is the well known and widely used technique of microarray analysis which has proven to be extremely useful for the identification of mRNA molecules that are differentially expressed in one tissue or cell type relative to another. In the course of our research using microarray analysis, we have identified approximately 200 gene transcripts that are present in human tumor cells at significantly higher levels than in corresponding normal human cells. To date, we have generated antibodies that bind to about 30 of the tumor antigen proteins expressed from these differentially expressed gene transcripts and have used these antibodies to quantitatively determine the level of production of these tumor antigen proteins in both human cancer cells and corresponding normal cells. We have then compared the levels of mRNA and protein in both the tumor and normal cells analyzed.
5. From the mRNA and protein expression analyses described in paragraph 4 above, we have observed that there is a strong correlation between changes in the level of mRNA present in any particular cell type and the level of protein

expressed from that mRNA in that cell type. In approximately 80% of our observations we have found that increases in the level of a particular mRNA correlates with changes in the level of protein expressed from that mRNA when human tumor cells are compared with their corresponding normal cells.

6. Based upon my own experience accumulated in more than 20 years of research, including the data discussed in paragraphs 4 and 5 above and my knowledge of the relevant scientific literature, it is my considered scientific opinion that for human genes, an increased level of mRNA in a tumor cell relative to a normal cell typically correlates to a similar increase in abundance of the encoded protein in the tumor cell relative to the normal cell. In fact, it remains a central dogma in molecular biology that increased mRNA levels are predictive of corresponding increased levels of the encoded protein. While there have been published reports of genes for which such a correlation does not exist, it is my opinion that such reports are exceptions to the commonly understood general rule that increased mRNA levels are predictive of corresponding increased levels of the encoded protein.

7. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: 5/07/04

By: Paul Polakis

Paul Polakis, Ph.D.

CURRICULUM VITAE

PAUL G. POLAKIS
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EDUCATION:

Ph.D., Biochemistry, Department of Biochemistry,
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PROFESSIONAL EXPERIENCE:

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Staff Scientist, Genentech, Inc
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1999- 2002

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Research Director
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1992- 1996

Senior Scientist, Project Leader, Onyx
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~~1991-1992~~

~~Senior Scientist, Chiron Corporation,
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1989-1991

Scientist, Cetus Corporation, Emeryville CA.

1987-1989

Postdoctoral Research Associate, Genentech,
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1985-1987

Postdoctoral Research Associate, Department
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1984-1985

Assistant Professor, Department of Chemistry,
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1980-1984

Graduate Research Assistant, Department of
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PUBLICATIONS:

1. Polakis, P. G. and Wilson, J. E. 1982 Purification of a Highly Bindable Rat Brain Hexokinase by High Performance Liquid Chromatography. **Biochem. Biophys. Res. Commun.** 107, 937-943.
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EXHIBIT B

	tumor mRNA	tumor IHC
UNQ2525	+	+
UNQ2378	+	+
UNQ972	+	-
UNQ97671	+	+
UNQ2964	+	+
UNQ323	+	+
UNQ1655	+	+
UNQ2333	+	+
UNQ9638	+	+
UNQ8209	+	+
UNQ6507	+	+
UNQ8196	+	+
UNQ9109	+	+
UNQ100	+	+
UNQ178	+	+
UNQ1477	+	+
UNQ1839	+	+
UNQ2079	+	+
UNQ8782	+	+
UNQ9646	+	-
UNQ111	+	+
UNQ3079	+	+
UNQ8175	+	+
UNQ9509	+	+
UNQ10978	+	-
UNQ2103	+	+
UNQ1563	+	+
UNQ16188	+	+
UNQ13589	+	+
UNQ1078	+	+
UNQ879	+	+

SECOND DECLARATION OF PAUL POLAKIS, Ph.D.

I, Paul Polakis, Ph.D., declare and say as follows:

1. I am currently employed by Genentech, Inc. where my job title is Staff Scientist.
2. Since joining Genentech in 1999, one of my primary responsibilities has been leading Genentech's Tumor Antigen Project, which is a large research project with a primary focus on identifying tumor cell markers that find use as targets for both the diagnosis and treatment of cancer in humans.
3. As I stated in my previous Declaration dated May 7, 2004 (attached as Exhibit A), my laboratory has been employing a variety of techniques, including microarray analysis, to identify genes which are differentially expressed in human tumor tissue relative to normal human tissue. The primary purpose of this research is to identify proteins that are abundantly expressed on certain human tumor tissue(s) and that are either (i) not expressed, or (ii) expressed at detectably lower levels, on normal tissue(s).
4. In the course of our research using microarray analysis, we have identified approximately 200 gene transcripts that are present in human tumor tissue at significantly higher levels than in normal human tissue. To date, we have successfully generated antibodies that bind to 31 of the tumor antigen proteins expressed from these differentially expressed gene transcripts and have used these antibodies to quantitatively determine the level of production of these tumor antigen proteins in both human tumor tissue and normal tissue. We have then quantitatively compared the levels of mRNA and protein in both the tumor and normal tissues analyzed. The results of these analyses are attached herewith as Exhibit B. In Exhibit B, "+" means that the mRNA or protein was detectably overexpressed in the tumor tissue relative to normal tissue and "-" means that no detectable overexpression was observed in the tumor tissue relative to normal tissue.
5. As shown in Exhibit B, of the 31 genes identified as being detectably overexpressed in human tumor tissue as compared to normal human tissue at the mRNA level, 28 of them (i.e., greater than 90%) are also detectably overexpressed in human tumor tissue as compared to normal human tissue at the protein level. As such, in the cases where we have been able to quantitatively measure both (i) mRNA and (ii) protein levels in both (i) tumor tissue and (ii) normal tissue, we have observed that in the vast majority of cases, there is a very strong correlation between increases in mRNA expression and increases in the level of protein encoded by that mRNA.

6. Based upon my own experience accumulated in more than 20 years of research, including the data discussed in paragraphs 4-5 above and my knowledge of the relevant scientific literature, it is my considered scientific opinion that for human genes, an increased level of mRNA in a tumor tissue relative to a normal tissue more often than not correlates to a similar increase in abundance of the encoded protein in the tumor tissue relative to the normal tissue. In fact, it remains a generally accepted working assumption in molecular biology that increased mRNA levels are more often than not predictive of elevated levels of the encoded protein. In fact, an entire industry focusing on the research and development of therapeutic antibodies to treat a variety of human diseases, such as cancer, operates on this working assumption.
7. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: 3-29-00

By: Paul Polakis

Paul Polakis, Ph.D.

EXHIBIT A

DECLARATION OF PAUL POLAKIS, Ph.D.

I, Paul Polakis, Ph.D., declare and say as follows:

1. I was awarded a Ph.D. by the Department of Biochemistry of the Michigan State University in 1984. My scientific Curriculum Vitae is attached to and forms part of this Declaration (Exhibit A).
2. I am currently employed by Genentech, Inc. where my job title is Staff Scientist. Since joining Genentech in 1999, one of my primary responsibilities has been leading Genentech's Tumor Antigen Project, which is a large research project with a primary focus on identifying tumor cell markers that find use as targets for both the diagnosis and treatment of cancer in humans.
3. As part of the Tumor Antigen Project, my laboratory has been analyzing differential expression of various genes in tumor cells relative to normal cells. The purpose of this research is to identify proteins that are abundantly expressed on certain tumor cells and that are either (i) not expressed, or (ii) expressed at lower levels, on corresponding normal cells. We call such differentially expressed proteins "tumor antigen proteins". When such a tumor antigen protein is identified, one can produce an antibody that recognizes and binds to that protein. Such an antibody finds use in the diagnosis of human cancer and may ultimately serve as an effective therapeutic in the treatment of human cancer.
4. In the course of the research conducted by Genentech's Tumor Antigen Project, we have employed a variety of scientific techniques for detecting and studying differential gene expression in human tumor cells relative to normal cells, at genomic DNA, mRNA and protein levels. An important example of one such technique is the well known and widely used technique of microarray analysis which has proven to be extremely useful for the identification of mRNA molecules that are differentially expressed in one tissue or cell type relative to another. In the course of our research using microarray analysis, we have identified approximately 200 gene transcripts that are present in human tumor cells at significantly higher levels than in corresponding normal human cells. To date, we have generated antibodies that bind to about 30 of the tumor antigen proteins expressed from these differentially expressed gene transcripts and have used these antibodies to quantitatively determine the level of production of these tumor antigen proteins in both human cancer cells and corresponding normal cells. We have then compared the levels of mRNA and protein in both the tumor and normal cells analyzed.
5. From the mRNA and protein expression analyses described in paragraph 4 above, we have observed that there is a strong correlation between changes in the level of mRNA present in any particular cell type and the level of protein

expressed from that mRNA in that cell type. In approximately 80% of our observations we have found that increases in the level of a particular mRNA correlates with changes in the level of protein expressed from that mRNA when human tumor cells are compared with their corresponding normal cells.

6. Based upon my own experience accumulated in more than 20 years of research, including the data discussed in paragraphs 4 and 5 above and my knowledge of the relevant scientific literature, it is my considered scientific opinion that for human genes, an increased level of mRNA in a tumor cell relative to a normal cell typically correlates to a similar increase in abundance of the encoded protein in the tumor cell relative to the normal cell. In fact, it remains a central dogma in molecular biology that increased mRNA levels are predictive of corresponding increased levels of the encoded protein. While there have been published reports of genes for which such a correlation does not exist, it is my opinion that such reports are exceptions to the commonly understood general rule that increased mRNA levels are predictive of corresponding increased levels of the encoded protein.

7. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: 5/07/04

By: Paul Polakis

Paul Polakis, Ph.D.

CURRICULUM VITAE

PAUL G. POLAKIS
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EDUCATION:

Ph.D., Biochemistry, Department of Biochemistry,
Michigan State University (1984)

B.S., Biology. College of Natural Science, Michigan State University (1977)

PROFESSIONAL EXPERIENCE:

2002-present	Staff Scientist, Genentech, Inc S. San Francisco, CA
1999- 2002	Senior Scientist, Genentech, Inc., S. San Francisco, CA
1997 -1999	Research Director Onyx Pharmaceuticals, Richmond, CA
1992- 1996	Senior Scientist, Project Leader, Onyx Pharmaceuticals, Richmond, CA
1991-1992	Senior Scientist, Chiron Corporation, Emeryville, CA.
1989-1991	Scientist, Cetus Corporation, Emeryville CA.
1987-1989	Postdoctoral Research Associate, Genentech, Inc., South San Francisco, CA.
1985-1987	Postdoctoral Research Associate, Department of Medicine, Duke University Medical Center, Durham, NC

1984-1985

Assistant Professor, Department of Chemistry,
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1980-1984

Graduate Research Assistant, Department of
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PUBLICATIONS:

1. Polakis, P. G. and Wilson, J. E. 1982 Purification of a Highly Bindable Rat Brain Hexokinase by High Performance Liquid Chromatography. **Biochem. Biophys. Res. Commun.** 107, 937-943.

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EXHIBIT B

	tumor mRNA	tumor IHC
UNQ2525	+	+
UNQ2378	+	+
UNQ972	+	-
UNQ97671	+	+
UNQ2964	+	+
UNQ323	+	+
UNQ1655	+	+
UNQ2333	+	+
UNQ9638	+	+
UNQ8209	+	+
UNQ6507	+	+
UNQ8196	+	+
UNQ9109	+	+
UNQ100	+	+
UNQ178	+	+
UNQ1477	+	+
UNQ1839	+	+
UNQ2079	+	+
UNQ8782	+	+
UNQ9646	+	-
UNQ111	+	+
UNQ3079	+	+
UNQ8175	+	+
UNQ9509	+	+
UNQ10978	+	-
UNQ2103	+	+
UNQ1563	+	+
UNQ16188	+	+
UNQ13589	+	+
UNQ1078	+	+
UNQ879	+	+

Overexpression of a DEAD Box Protein (DDX1) in Neuroblastoma and Retinoblastoma Cell Lines*

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The DEAD box gene, *DDX1*, is a putative RNA helicase that is co-amplified with *MYCN* in a subset of retinoblastoma (RB) and neuroblastoma (NB) tumors and cell lines. Although gene amplification usually involves hundreds to thousands of kilobase pairs of DNA, a number of studies suggest that co-amplified genes are only overexpressed if they provide a selective advantage to the cells in which they are amplified. Here, we further characterize *DDX1* by identifying its putative transcription and translation initiation sites. We analyze *DDX1* protein levels in *MYCN/DDX1*-amplified NB and RB cell lines using polyclonal antibodies specific to *DDX1* and show that there is a good correlation with *DDX1* gene copy number, *DDX1* transcript levels, and *DDX1* protein levels in all cell lines studied. *DDX1* protein is found in both the nucleus and cytoplasm of *DDX1*-amplified lines but is localized primarily to the nucleus of nonamplified cells. Our results indicate that *DDX1* may be involved in either the formation or progression of a subset of NB and RB tumors and suggest that *DDX1* normally plays a role in the metabolism of RNAs located in the nucleus of the cell.

DEAD box proteins are a family of putative RNA helicases that are characterized by eight conserved amino acid motifs, one of which is the ATP hydrolysis motif containing the core amino acid sequence DEAD (Asp-Glu-Ala-Asp) (1–3). Over 40 members of the DEAD box family have been isolated from a variety of organisms including bacteria, yeast, insects, amphibians, mammals, and plants. The prototypic DEAD box protein is the translation initiation factor, eukaryotic initiation factor 4A, which, when combined with eukaryotic initiation factor 4B, unwinds double-stranded RNA (4). Other DEAD box proteins, such as p68, Vasa, and An3, can effectively and independently destabilize/unwind short RNA duplexes *in vitro* (5–7). Although some DEAD box proteins play general roles in cellular processes such as translation initiation (eukaryotic initiation factor 4A (4)), RNA splicing (PRP5, PRP28, and SPP81 in yeast (8–10)), and ribosomal assembly (SrmB in *Escherichia coli* (11)), the function of most DEAD box proteins remains unknown. Many of the DEAD box proteins found in higher eukaryotes are tissue- or stage-specific. For example, *PL10* mRNA is expressed only in the male germ line, and its product

has been proposed to have a specific role in translational regulation during spermatogenesis (12). Vasa and ME31B are maternal proteins that may be involved in embryogenesis (13, 14). p68, found in dividing cells (15), is believed to be required for the formation of nucleoli and may also have a function in the regulation of cell growth and division (16, 17). Other DEAD box proteins are implicated in RNA degradation, mRNA stability, and RNA editing (18–20).

The human DEAD box protein gene *DDX1*¹ was identified by differential screening of a cDNA library enriched in transcripts present in the two RB cell lines Y79 and RB522A (21). The longest *DDX1* cDNA insert isolated from this library was 2.4 kb with an open reading frame from position 1 to 2201. All eight conserved motifs characteristic of DEAD box proteins are found in the predicted amino acid sequence of *DDX1* as well as a region with homology to the heterogeneous nuclear ribonucleoprotein U, a protein believed to participate in the processing of heterogeneous nuclear RNA to mRNA (22, 23). The region of homology to heterogeneous nuclear ribonucleoprotein U spans 128 amino acids and is located between the first two conserved DEAD box protein motifs, 1a and 1b.

The proto-oncogene *MYCN* encodes a member of the MYC family of transcription factors that bind to an E box element (CACGTG) when dimerized with the MAX protein (24, 25). The *MYCN* gene is amplified and overexpressed in approximately one-third of all NB tumors (26, 27). Amplification of *MYCN* is associated with rapid tumor progression and a poor clinical prognosis (26, 27). *MYCN* overexpression is usually achieved by increasing gene copy number rather than by up-regulating basal expression of *MYCN* (27, 28). Because gene amplification involves hundreds to thousands of kilobase pairs of contiguous DNA (29–32), it is possible that co-amplification of a gene located in proximity to *MYCN* may contribute to the poor clinical prognosis of *MYCN*-amplified tumors. The *DDX1* gene maps to the same chromosomal band as *MYCN*, 2p24, and is located ~400 kb telomeric to the *MYCN* gene (33–36). All four *MYCN*-amplified RB tumor cell lines tested to date are amplified for *DDX1* (21),² while approximately two-thirds of NB cell lines and 38–68% of NB tumors are co-amplified for both genes (37–39). George *et al.* (39) found a significant decrease in the mean disease-free survival of patients with *DDX1/MYCN*-amplified NB tumors compared with *MYCN*-amplified tumors. Similarly, Squire *et al.* (38) observed a trend toward a worse clinical prognosis when both genes were amplified in the tumors of NB patients. To date, there have been no reports of a

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The nucleotide sequence(s) reported in this paper has been submitted to the GenBank™/EBI Data Bank with accession number(s) X70649.

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¹ The abbreviations used are: *DDX1*, DEAD box 1; NB, neuroblastoma; RB, retinoblastoma; RACE, rapid amplification of cDNA ends; PAGE, polyacrylamide gel electrophoresis; nt, nucleotide(s); MOPS, 4-morpholinepropanesulfonic acid; bp, base pair(s); kb, kilobase(s) or kilobase pair(s).

² R. Godbout, unpublished results.

tumor amplified only for *DDX1*, and the role that this gene plays in cancer formation and progression is not known.

Because of the high rate of rearrangements in amplified DNA (31, 40), it is unlikely that a gene located ~400 kb from the *MYCN* gene will be consistently amplified as an intact unit unless its product provides a growth advantage to the cell. Based on Southern blot analysis, the *DDX1* gene extends over more than 30 kb, and there are no gross rearrangements of this gene in *DDX1*-amplified tumors (21, 38). Furthermore, there is a good correlation between *DDX1* transcript levels and gene copy number in the tumors analyzed to date. However, we need to show that *DDX1* protein is overexpressed in *DDX1*-amplified tumors if we are to entertain the possibility that this protein plays a role in the tumorigenic process. Here, we isolate and characterize the 5'-end of *DDX1* mRNA and extend the *DDX1* cDNA sequence by ~300 nt. We identify the predicted initiation codon of *DDX1* and generate antisera that specifically recognize *DDX1* protein. We analyze levels of *DDX1* protein in both *DDX1*-amplified and nonamplified RB and NB tumors and study the subcellular location of this protein in the cell.

MATERIALS AND METHODS

Library Screening—A human fetal brain cDNA library (Stratagene) was screened using a 320-bp DNA fragment from the 5'-end of the 2.4-kb *DDX1* cDNA previously described (23). Phagemids containing positive inserts were excised from λ ZAP II following the supplier's directions. The ends of the cDNA inserts were sequenced using the dideoxynucleotide chain termination method with T7 DNA polymerase (Amersham Pharmacia Biotech).

A human placenta genomic library (CLONTECH) was screened with the 5'-end of *DDX1* cDNA. Positive plaques were purified, and the genomic DNA was analyzed using restriction enzymes and Southern blotting. *EcoRI*-digested DNA fragments from these clones were subcloned into pBluescript and digested with exonuclease III and mung bean nuclease to obtain sequentially deleted clones. The exon/intron map of the 5' portion of the *DDX1* gene was obtained by comparing the sequence of *DDX1* cDNA with that of the genomic DNA.

Rapid Amplification of cDNA Ends (RACE)—We used the AmpliFINDER RACE kit (CLONTECH) to extend the 5'-end of *DDX1* cDNA. Briefly, two μ g of poly(A)⁺ RNA isolated from RB522A was reverse transcribed at 52 °C using either primer P1 or P3 (Fig. 1A). The RNA template was hydrolyzed, and excess primer was removed. A single-stranded AmpliFINDER anchor containing an *EcoRI* site was ligated to the 3'-end of the cDNA using T4 RNA ligase. The cDNA was amplified using either primer P2 or P4 (Fig. 1A) and AmpliFINDER anchor primer. RACE products were cloned into pBluescript.

Primer Extension—Poly(A)⁺ RNAs were isolated from RB and NB cell lines as described previously (21, 38). The 21-nt primers 5'-TTCGT-TCTGGG-CACCATGTGT-3' (primer P4 in Fig. 1A) and 5'-TGGGAC-CTAGGGCTTCTGGAC-3' (primer P3 in Fig. 1A) were end-labeled with [γ -³²P]ATP (3000 Ci/mmol; Mandel Scientific) and T4 polynucleotide kinase. Each of the labeled primers was annealed to 2 μ g of poly(A)⁺ RNA at 45 °C for 90 min, and the cDNA was extended at 42 °C for 60 min using avian myeloblastosis virus reverse transcriptase (Promega). The primer extension products were heat-denatured and run on a 8% polyacrylamide gel containing 7 M urea in 1 \times TBE buffer. A G + A sequencing ladder served as the size standard.

S1 Nuclease Protection Assay—The S1 nuclease protection assay to map the transcription initiation site of *DDX1* was performed as described by Favalaro *et al.* (41). The DNA probe was prepared by digesting genomic DNA spanning the upstream region of *DDX1* and exon 1 with *AvaI*, labeling the ends with [γ -³²P]ATP (3000 Ci/mmol) and polynucleotide kinase, and removing the label from one of the ends by digesting the DNA with *SphI* (Fig. 4). The RNA samples were resuspended in a hybridization mixture containing 80% formamide, 40 mM PIPES, 400 mM NaCl, 1 mM EDTA, and the heat-denatured *SphI*-*AvaI* probe labeled at the *AvaI* site. The samples were incubated at 45 °C for 16 h and digested with 3000 units/ml S1 nuclease (Boehringer Mannheim) for 60 min at 37 °C. The samples were precipitated with ethanol; resuspended in 80% formaldehyde, TBE buffer, 0.1% bromophenol blue, xylene cyanol; denatured at 90 °C for 2 min; and electrophoresed in a 7 M urea, 8% polyacrylamide gel in TBE buffer.

Northern and Southern Blot Analysis—Poly(A)⁺ RNAs were isolated from RB and NB cell lines as described previously (21, 38). Two μ g of

poly(A)⁺ RNA/lane were electrophoresed in a 6% formaldehyde, 1.5% agarose gel in MOPS buffer (20 mM MOPS, 5 mM sodium acetate, 1 mM EDTA, pH 7.0) and transferred to nitrocellulose filter in 3 M sodium chloride, 0.3 M sodium citrate. The filters were hybridized to the following DNA probes, ³²P-labeled by nick translation: (i) a 1.6-kb *EcoRI* insert from *DDX1* cDNA clone 1042 (21), (ii) a 260-bp cDNA fragment spanning the 3'-end of *DDX1* exon 1 as well as exons 2 and 3, (iii) a 160-bp fragment derived from the 5'-end of *DDX1* exon 1, and (iv) α -actin cDNA to control for lane to lane variation in RNA levels. Filters were hybridized and washed under high stringency. Southern blot analysis was as described previously (21).

Preparation of Anti-DDX1 Antiserum—To prepare antiserum to the C terminus of the *DDX1* protein, we inserted a 1.8-kb *EcoRI* fragment from bp 848 to 2668 of *DDX1* cDNA (Fig. 1B) into *EcoRI*-digested pMAL-c2 expression vector (New England Biolabs). DH5 α cells transformed with this vector were grown to mid-log phase and induced with 0.1 mM isopropyl-1-thio- β -D-thiogalactoside. The cells were harvested 3–4 h postinduction and lysed by sonication. Soluble maltose binding protein-DDX1 fusion protein was affinity-purified using amylose resin, and the maltose-binding protein was cleaved with factor Xa. The *DDX1* protein was purified on a SDS-PAGE gel, electroeluted, and concentrated. Approximately 100 μ g of protein was injected into rabbits at 4–6-week intervals. For the initial injection, the protein was dispersed in complete Freund's adjuvant (Sigma), while subsequent injections were prepared in Freund's incomplete adjuvant. Blood was collected from each rabbit 10 days after injection, and the specificity of the antiserum was tested using cell extracts from RB522A. To prepare antiserum to the N terminus of *DDX1* protein, a *DDX1* cDNA fragment from bp 268 to 851 (Fig. 1B) was inserted into pGEX-4T2 (Amersham Pharmacia Biotech). The recombinant protein produced from this construct contains the first 186 amino acids of the predicted *DDX1* sequence. Soluble glutathione S-transferase-DDX1 fusion protein was purified with glutathione-Sepharose 4B (Amersham Pharmacia Biotech). The glutathione S-transferase component of the fusion protein was cleaved with thrombin.

Subcellular Fractionations and Western Blot Analysis—We used two different procedures for subcellular fractionations. First, we isolated nuclear and S100 (soluble cytoplasmic) fractions from RB522A, IMR-32, Y79, RB(E)-2, HeLa, and HL60 using the procedure of Dignam (42). On average, we obtained 5–6 times more protein in the cytosolic fractions than in the nuclear fractions. Second, 10⁸ RB522A cells were lysed and fractionated into S4 (soluble cytoplasmic components), P2 (heavy mitochondria, plasma membrane fragments), P3 (mitochondria, lysosomes, peroxisomes, and Golgi membranes), and P4 fractions (membrane vesicles from rough and smooth endoplasmic reticulum, Golgi, and plasma membrane) by differential centrifugation (43). We obtained 8 mg of protein in the S4 fraction, 1 mg in P2, 0.5 mg in P3, and 2 mg in P4 fraction. The procedures related to the immunoelectron microscopy have been previously described (44).

For Western blot analysis, proteins were electrophoresed in polyacrylamide-SDS gels and electroblotted onto nitrocellulose using the standard protocol for protein transfer described by Schleicher and Schuell. The filters were incubated with a 1:5000 dilution of *DDX1* antiserum, a 1:200 dilution of anti-MYCIN monoclonal antibody (Boehringer Mannheim), or a 1:200 dilution of anti-actin (Santa Cruz Biotechnology, Inc., Santa Cruz, CA). For the colorimetric analysis, antigen-antibody interactions were visualized using either alkaline phosphatase-linked goat anti-rabbit IgG (for *DDX1*) or goat anti-mouse IgG (for MYCIN) at a 1:3000 dilution. For the ECL Western blotting analysis (Amersham Pharmacia Biotech), we used a 1:100,000 dilution of peroxidase-linked secondary anti-rabbit IgG antibody (for *DDX1*) or secondary anti-goat IgG antibody (Jackson ImmunoResearch Laboratories).

RESULTS

Identification of the 5'-End of the *DDX1* Transcript—We have previously reported the sequence of *DDX1* cDNA isolated from an RB cDNA library (21, 23). This 2.4-kb *DDX1* cDNA contains an open reading frame spanning positions 1–2201 with a methionine encoded by the first three nucleotides (Fig. 1A). There is a polyadenylation signal and poly(A) tail in the 3'-untranslated region, indicating that the sequence is complete at the 3'-end. Manohar *et al.* (37) have also isolated *DDX1* cDNA from the NB cell line LA-N-5. Their cDNA extended the 5'-end of our sequence by 42 bp and included an additional in frame methionine (*double underlined* in Fig. 1A). The possibil-

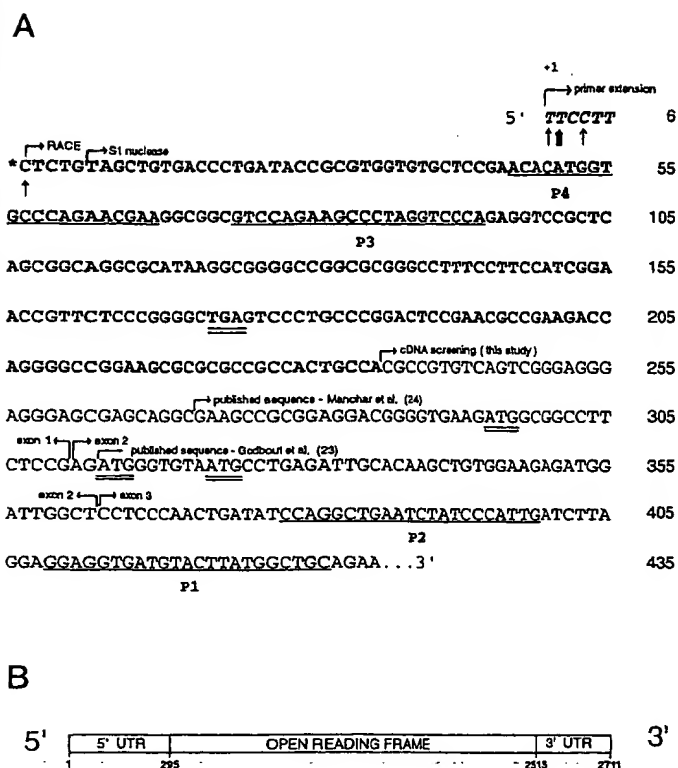


FIG. 1. Partial sequence and structure of *DDX1* cDNA. A, the sequence of the 5'-end of *DDX1* cDNA. The sequence in **boldface type** starting at the asterisk was obtained using the RACE strategy. The additional 6 bp in *italic boldface type* at the 5'-end of the cDNA are predicted based on the known *DDX1* genomic sequence and primer extension analysis. P1, P2, P3, and P4 are primers used in the RACE experiments (the complementary sequence was used in each case). Primers P3 and P4 were also used for the primer extension analysis. Three in frame methionine codons are indicated by the **double underline**. An in frame stop codon is indicated by the **boldface double underline**. The three major transcription initiation sites identified by primer extension are indicated by the **single arrows**, while a minor site is represented by the **broad arrow**. The predicted *DDX1* transcription initiation sites obtained by RACE, S1 nuclease, and primer extension are indicated as well as the 5'-ends of *DDX1* cDNA sequences obtained by screening cDNA libraries. The sequences transcribed from exons 1, 2, and 3 are also shown. B, the structure of the 2711-bp *DDX1* cDNA is shown with an open reading frame from position 295 to 2515.

ity of additional in frame methionines located further upstream could not be excluded, because there were no predicted stop codons in the upstream region of the cDNA.

Northern blot analysis indicated a *DDX1* transcript size of ~2800 nt, suggesting that the *DDX1* cDNAs isolated to date were lacking ~300–350 bp of 5' sequence. We have used different approaches to identify the transcription start site of *DDX1*. First, we exhaustively screened a commercial fetal brain cDNA library with the 5'-end of *DDX1* cDNA. Although numerous clones were analyzed, only one extended the sequence (by 35 bp) beyond that published by Manohar *et al.* (37) (Fig. 1A).

We next used the RACE procedure in an attempt to isolate additional 5' sequence. The nested primers used to amplify the 5'-end of the *DDX1* transcript are labeled as primers P1 and P2 in Fig. 1A and are located downstream of the three in frame methionines (**double underline** in Fig. 1A). Poly(A)⁺ RNA from RB522A was reverse transcribed at 52 °C using primer P1, and the reverse transcribed cDNA was amplified using the nested primer P2 and the 5'-RACE primer. Using this approach, we generated a product that was 230 bp longer than any of the cDNAs obtained by screening libraries (Fig. 1A). Sequencing of this 230-bp cDNA revealed an in frame stop codon (**boldface double underline** in Fig. 1A) located 123 bp

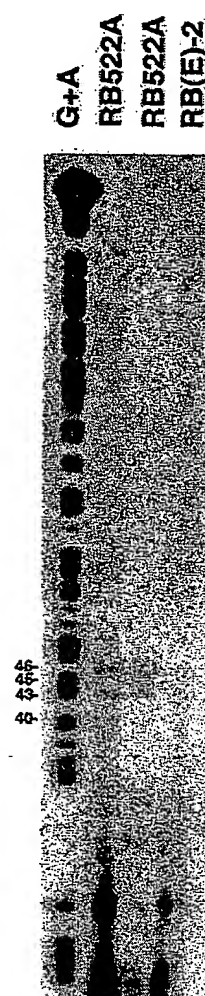


FIG. 2. Identification of the 5'-end of the *DDX1* transcript by primer extension. Radioactively labeled primer P4 was annealed to 2 µg of poly(A)⁺ RNA from RB522A (lane 1), 1 µg of poly(A)⁺ RNA from RB522A (lane 2), and 2 µg of poly(A)⁺ RNA from RB(E)-2 cells (lane 3), and extended using reverse transcriptase. The products were run on an 8% denaturing polyacrylamide gel with a G + A sequencing ladder as size marker. The primer extension products are indicated on the left. The sizes of the products (in nt) are presented as the distance from primer P4.

upstream of the predicted translation initiation site. We then prepared primers P3 and P4, located near the 5'-end of the RACE cDNA (Fig. 1A) and repeated the RACE procedure to see if additional 5' sequences could be obtained. The resulting RACE products did not extend the *DDX1* cDNA sequence further.

The location of the *DDX1* transcription initiation site was verified by primer extension. Poly(A)⁺ RNA was prepared from the following two cell lines: *DDX1*-amplified RB cell line RB522A and a nonamplified RB cell line RB(E)-2. RB522A has elevated levels of *DDX1* mRNA, while RB(E)-2 has at least 20-fold lower levels of this transcript. Three products of 40, 43, and 46 nt (with a weak signal at 45 nt) were detected in RB522A using primer P4 (Figs. 1A and 2). The 40-nt product corresponded exactly with the 5'-end of the RACE-derived cDNA while the 43- and 46-nt products extended the predicted size of the *DDX1* transcript by 3 and 6 nt, respectively. None of these products were observed in RB(E)-2. Bands of identical sizes to those obtained with RB522A mRNA were also observed in the *DDX1*-amplified NB cell line BE(2)-C but not in the *DDX1*-amplified NB cell line IMR-32 (data not shown). The same predicted *DDX1* transcription initiation site was identified with primer P3 except that the bands were of weaker

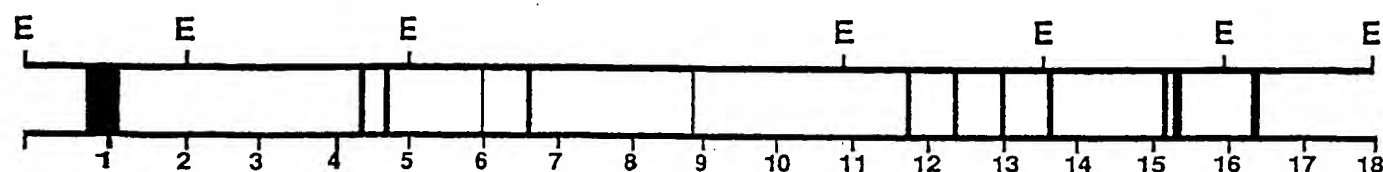


FIG. 3. Genomic map of the 5'-end of *DDX1*. The exons are represented by the black boxes, and distances are in kilobase pairs. The locations of *EcoRI* (E) sites are indicated.

intensity (data not shown). We have designated the transcription start site identified by primer extension as +1 (Fig. 1A).

The sequence of the 6 nt extending beyond the RACE cDNA was obtained by comparison of the cDNA sequence with that of *DDX1* genomic DNA. Bacteriophages containing *DDX1* genomic DNA were isolated by screening a human placenta library with 5' *DDX1* cDNA. Eighteen kb of DNA were sequenced from two bacteriophages with overlapping *DDX1* genomic DNA. Thirteen exons were identified within this 18-kb region (Fig. 3) corresponding to cDNA sequences from position 1 to 1249. The 310-bp exon 1 was by far the longest of the 13 exons sequenced, corresponding to the entire 5'-untranslated region of *DDX1* as well as the first in frame methionine. The sequences transcribed from exons 1, 2, and 3 are indicated in Fig. 1A.

Knowledge of the genomic structure of *DDX1* allowed us to use the S1 protection assay, a technique that is independent of reverse transcriptase, to further define the 5'-end of the *DDX1* transcript. Poly(A)⁺ RNAs from six *DDX1*-amplified lines (RB lines: Y79 and RB522A; NB lines: BE(2)-C, IMR-32, LA-N-1, and LA-N-5) and six nonamplified lines (RB lines: RB(E)-2 and RB412; NB lines: GOTO, NB-1, NUB-7, and SK-N-MC) were hybridized to a DNA probe that extended from position -745 in the 5'-flanking *DDX1* DNA to position +164 in exon 1. This DNA probe was labeled at position +164 as indicated in Fig. 4. Nonhybridized DNA was digested with S1 nuclease, and the sizes of the protected fragments were analyzed on a denaturing polyacrylamide gel. Bands of 150–153 nt were observed in lane 2 (RB522A), lane 5 (BE(2)-C), and lane 8 (LA-N-1) with bands of much weaker intensity in lane 7 (IMR-32) (Fig. 4). Specific bands were not detected in either *DDX1*-amplified Y79 and LA-N-5 or the nonamplified lines. Although the sizes of the S1 protected bands in RB522A, BE(2)-C, and LA-N-1 were 5 and 11 nt shorter than predicted based on RACE and primer extensions, respectively, there was general agreement with all three techniques regarding the location of the *DDX1* transcription initiation site (Fig. 1A). The smaller S1 nuclease protected products could have arisen as the result of S1 digestion of the 5'-end of the RNA:DNA heteroduplex because of its relatively high rU:dA content (45).

Identification of the same transcription initiation site in three *DDX1*-amplified lines suggests that this represents the bona fide start site of *DDX1* transcription. However, it was not clear why this start site was either very weak or not detected in three other amplified lines. To determine whether the 5'-end of exon 1 is transcribed in all *DDX1*-amplified lines, we carried out a direct analysis of the 5'-end of the *DDX1* transcript by Northern blotting. Two probes were used for this analysis: the 5' probe contained a 160-bp fragment from bp 1 to 160 (5'-half of exon 1), and the 3' probe contained a 260-bp fragment from bp 160 to 420 (3'-half of exon 1 as well as exons 2 and 3) (Fig. 1A). With the 3' probe, we obtained bands of similar size and intensity in four *DDX1*-amplified lines (RB522A, BE(2)-C, IMR-32, and LA-N-5). Band intensity was somewhat weaker in Y79 and stronger in LA-N-1 in comparison with the other lines (Fig. 5). No signal was detected in the non-*DDX1*-amplified line RB412. With the 5' probe, a relatively strong signal was observed in RB522A, BE(2)-C, and LA-N-1, while a considerably weaker

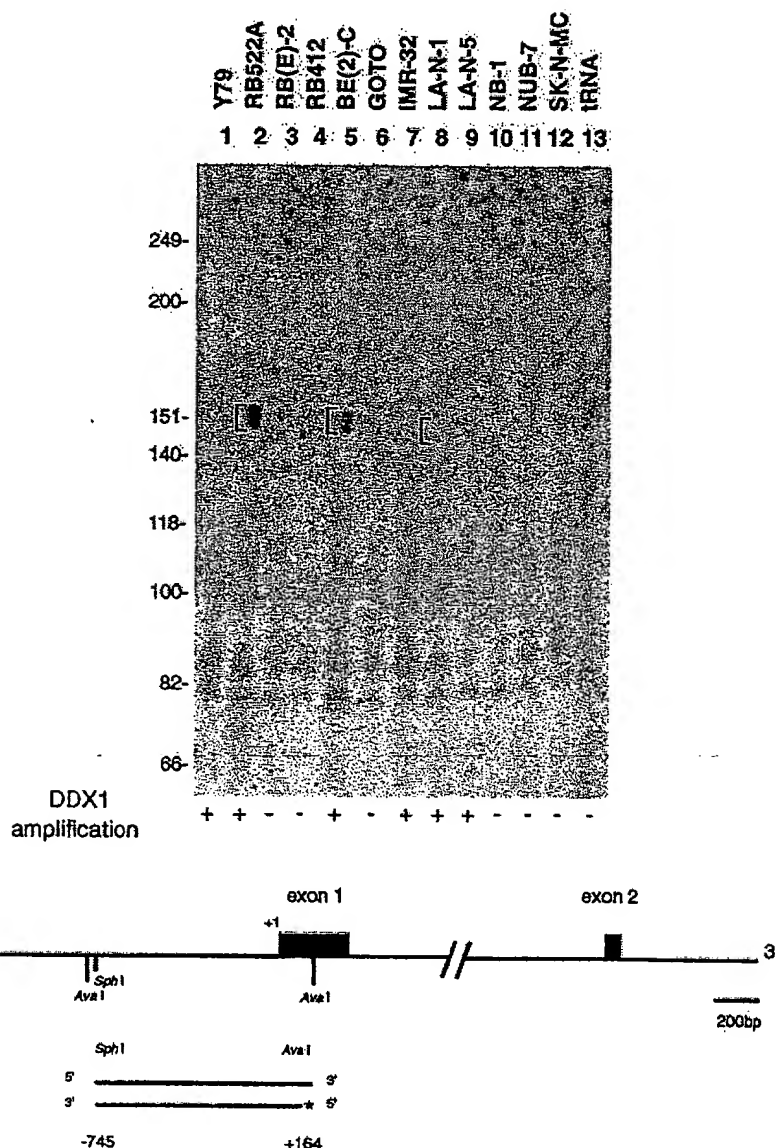
but readily apparent signal was detected in Y79, IMR-32, and LA-N-5. The signal obtained with actin indicates that, with the exception of LA-N-1, similar amounts of RNA were loaded in each lane and that the RNA was not degraded. These results indicate that at least a portion of the 160-bp 5'-end of exon 1 is transcribed in all *DDX1*-amplified lines.

Based on primer extension, S1 nuclease protection assay, Northern blot analysis and the sequencing of the RACE products, we conclude that the *DDX1* transcript is 2.7 kb with an open reading frame spanning nucleotides 295–2515 encoding a predicted protein of 740 amino acids with an estimated molecular weight of 82.4 (Fig. 1B). An in frame stop codon is located 123 nt upstream of the predicted translation initiation site, at positions 172–174. The first in frame methionine following the stop codon is in agreement with the Kozak consensus sequence (46). Furthermore, the predicted start methionine codon for human *DDX1* corresponds perfectly with that of *Drosophila* *DDX1* (47). A stop codon is located 15 nt upstream of the initiation codon in *Drosophila* *DDX1*.

Analysis of *DDX1* Protein Levels in Neuroblastoma and Retinoblastoma—We and others have previously shown that there is a good correlation between gene copy number and RNA levels in *DDX1*-amplified RB and NB cell lines (37, 38). To determine whether the correlation extends to *DDX1* protein levels, we prepared antiserum to two nonoverlapping recombinant *DDX1* proteins. First, we prepared a C terminus recombinant protein construct by inserting a 1.8-kb *EcoRI* fragment from bp 848 to 2668 (amino acids 185–740) (Fig. 1B) into the pMAL-c2 expression vector. Recombinant protein expression was induced with isopropyl-1-thio- β -D-thiogalactoside, and the 110-kDa maltose-binding protein-*DDX1* fusion product was purified by affinity chromatography using amylose resin, followed by electrophoresis on a SDS-PAGE gel after cleaving the maltose-binding protein fusion partner with factor Xa. Second, we prepared an N terminus construct by ligating a DNA fragment from bp 268 to 851 (amino acids 1–186) into pGEX-4T2. The 50-kDa glutathione S-transferase-*DDX1* fusion protein was purified by affinity chromatography on a glutathione column. This N terminus fusion protein contains only the first of the eight conserved motifs found in all DEAD box proteins, while the C terminus fusion protein includes the remaining seven motifs.

We measured *DDX1* protein levels in total cell extracts of three RB and 10 NB cell lines. Using antiserum to the N terminus fusion protein, we observed a strong signal in all *DDX1*-amplified cell lines: the RB cell lines Y79 (lane 1) and RB522A (lane 2) and the NB cell lines BE(2)-C (lane 4), IMR-32 (lane 6), LA-N-1 (lane 8), and LA-N-5 (lane 9) (Fig. 6). Two bands were observed in the majority of extracts. Of the amplified lines, Y79 produced the weakest signal, with the most intense signal observed in LA-N-1. There was an excellent correlation with *DDX1* protein and mRNA levels in these cell lines, with lower levels of *DDX1* mRNA observed in Y79 and higher levels in LA-N-1 (Fig. 7A). As shown in Fig. 7B, this correlation extended to *DDX1* gene copy number. No gross DNA rearrangements were seen in the *DDX1*-amplified lines; however, three small bands of altered size were observed in the RB412 lane. Although the nature of the DNA alteration is not known, it is noteworthy that *DDX1* transcript levels in RB412

FIG. 4. S1 nuclease mapping of the 5'-end of the *DDX1* transcript. Two μ g of poly(A)⁺ RNA from four RB lines (*DDX1*-amplified Y79 and RB522A and nonamplified RB(E)-2 and RB412), eight NB lines (*DDX1*-amplified BE(2)-C, IMR-32, LA-N-1, and LA-N-5 and nonamplified GOTO, NB-1, NUB-7, and SK-N-MC), and tRNA as a negative control were hybridized to a *Sph*I-*Ava*I fragment labeled at the *Ava*I site with [γ -³²P]ATP and polynucleotide kinase. Bands of 150–153 nt are shown in lanes 2 (RB522A), 5 (BE(2)-C), and 8 (LA-N-1) with much weaker bands in lane 7 (IMR-32). A map of the probe indicating the transcription initiation site identified by primer extension (+1), the labeling site (*), and exons 1 and 2, is shown at the bottom.



are extremely low (Fig. 7A) and that the top *DDX1* protein band in RB412 cell extracts is smaller in size than the top band from the other cell extracts (Fig. 6).

Two *DDX1* protein bands were present in most of the lanes in Fig. 6. The same two bands were detected with antiserum to the C terminus of the *DDX1* protein, as well as a third band at ~60 kDa (data not shown). There was no variation in the intensity of the 60-kDa band in *DDX1*-amplified and nonamplified cell extracts. The 60-kDa band probably represents another member of the DEAD box protein family, because the C terminus *DDX1* protein used to prepare this antiserum contained seven of the eight conserved motifs found in all DEAD box proteins. To obtain an estimate of the size of the two *DDX1* bands, we ran cellular extracts from RB522A on a 7% SDS-PAGE gel with the BenchMark protein ladder (Life Technologies, Inc.). The size of the *DDX1* protein was determined using the Alpha Imager 2000 documentation and analysis system for molecular weight calculation. Based on this analysis, the estimated molecular mass of the top band is 89.5 kDa, while that of the bottom band is 83.5 kDa. The 84-kDa band may represent the unmodified product encoded by the *DDX1* transcript (capable of encoding a protein with a predicted molecular mass of 82.4 kDa), while the top band may represent post-translational modification of *DDX1* protein (e.g. phosphorylation). Another possibility is that the top band represents intact *DDX1*

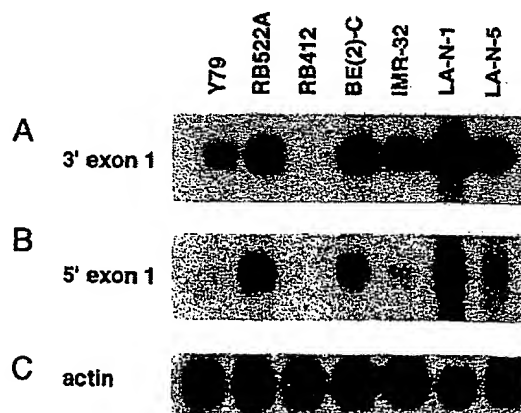


FIG. 5. Northern blot analysis of the 5'-end of the *DDX1* transcript. Two μ g of poly(A)⁺ RNA isolated from *DDX1*-amplified Y79, RB522A, BE(2)-C, IMR-32, LA-N-1, and LA-N-5 and nonamplified RB412 were electrophoresed in a 1.5% agarose-formaldehyde gel. The RNA was transferred to a nitrocellulose filter and sequentially hybridized with a 260-bp fragment from *DDX1* cDNA from bp +160 to +420 (3'-end of exon 1 as well as exons 2 and 3) (A), a 160-bp fragment from *DDX1* cDNA from bp +1 to +160 (5'-end of exon 1) (B), and actin cDNA (C). The DNA was labeled with [³²P]dCTP by nick translation. The blots were hybridized and washed under high stringency.

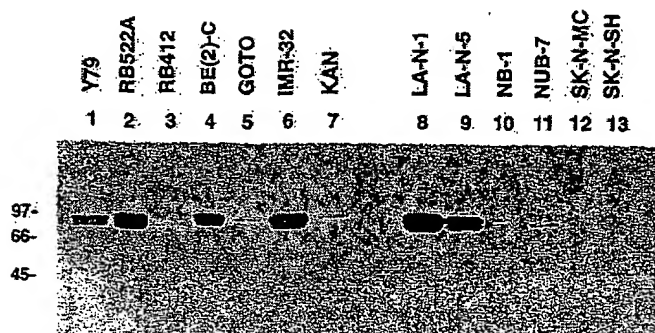


FIG. 6. DDX1 protein expression in RB and NB cell lines. Western blots were prepared using total cellular extracts from three RB (Y79, RB522A, and RB412) and 10 NB cell lines (BE(2)-C, GOTO, IMR-32, KAN, LA-N-1, LA-N-5, NB-1, NUB-7, SK-N-MC, and SK-N-SH). The lines that are amplified for the *DDX1* gene are Y79, RB522A, BE(2)-C, IMR-32, LA-N-1, and LA-N-5. Twenty μ g of protein were loaded in each lane and electrophoresed in a 10% SDS-PAGE gel. DDX1 was detected using a 1:5000 dilution of the antiserum to the amino terminus of DDX1 protein. Size markers in kilodaltons are indicated on the side.

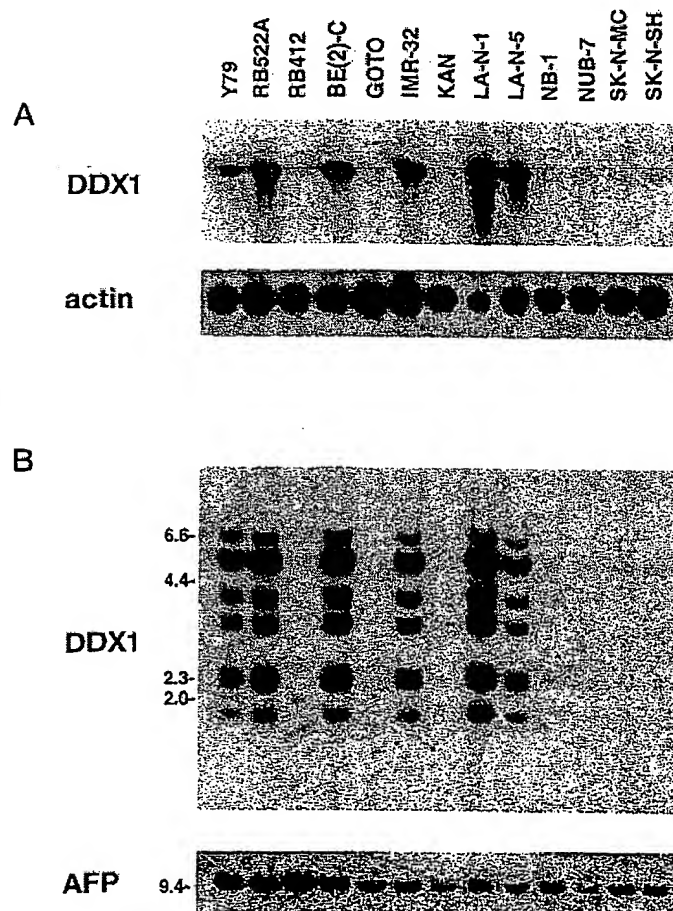


FIG. 7. Northern and Southern blot analyses of *DDX1* in RB and NB cell lines. A, 2 μ g of poly(A)⁺ RNA were loaded in each lane, electrophoresed in a 1.5% agarose-formaldehyde gel, and transferred to a nitrocellulose filter. The filter was first hybridized to a ³²P-labeled 1.6-kb *DDX1* cDNA (clone 1042) (21), stripped, and rehybridized to actin DNA. B, 10 μ g of genomic DNA from each of the indicated cell lines were digested with *Eco*RI, electrophoresed in a 1% agarose gel, and transferred to a nitrocellulose filter. The filter was hybridized to ³²P-labeled clone 1042 *DDX1* cDNA, stripped, and rehybridized with labeled α -fetoprotein cDNA. Markers (in kilobase pairs) are indicated on the side.

and the lower band is a specific truncated or degradation product of DDX1. Yet a third possibility is that the two bands represent the products of differentially spliced transcripts or

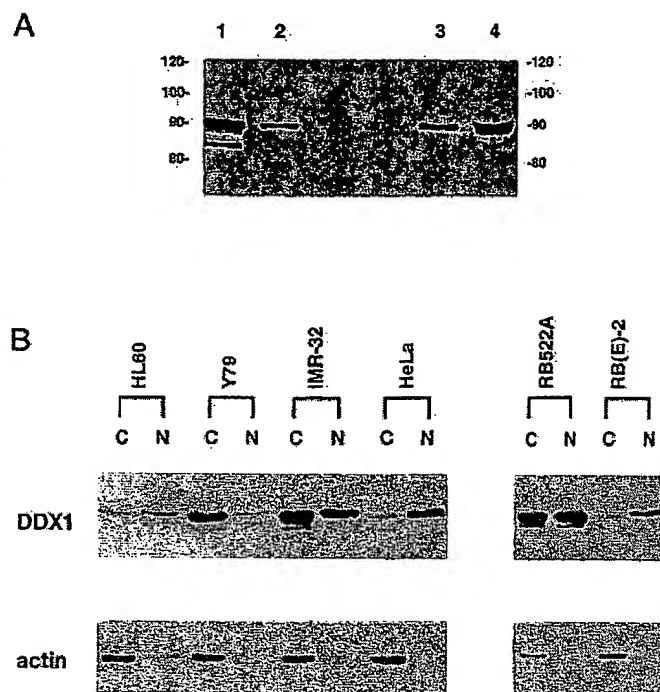


FIG. 8. Distribution of DDX1 in the nucleus and cytoplasm. A, cytosolic and nuclear extracts were prepared from RB522A and electrophoresed in a 7% SDS-PAGE gel. Cytosolic extracts were loaded in lanes 1 (20 μ g of protein) and 2 (10 μ g), while nuclear extracts were loaded in lanes 3 (10 μ g) and 4 (20 μ g). DDX1 was visualized using a 1:5000 dilution of the antiserum to the N terminus. The BenchMark protein ladder size markers (kilodaltons) are indicated on the left. B, cytosolic and nuclear extracts were prepared from HL60, Y79, IMR-32, HeLa, RB522A, and RB(E)-2 and electrophoresed in an 8% SDS-PAGE gel. Twenty μ g of proteins were loaded in each lane marked C (cytosolic) and N (nuclear). DDX1 was visualized using a 1:5000 dilution of the antiserum to the N terminus. Actin levels were analyzed using a 1:200 dilution of anti-actin antibody (Santa Cruz Biotechnology).

different translation initiation sites. However, the lack of any obvious differences in *DDX1* transcript sizes in the three RB and 10 NB lines analyzed in Fig. 7A does not support the latter possibility (e.g. compare the *DDX1* transcript size in NUB-7 (which produces the lower DDX1 protein band) and in NB-1 (which produces the higher DDX1 protein band)).

Subcellular Localization of DDX1 Protein—DEAD box proteins have been implicated in a variety of cellular functions including RNA splicing in the nucleus, translation initiation in the cytoplasm, and ribosome assembly in the nucleolus. To obtain an indication of the possible role of DDX1, we studied its subcellular location. Nuclear and cytosolic extracts were prepared from *DDX1*-amplified RB522A and run on a 7% SDS-PAGE gel. Although there was more DDX1 protein in the cytosol than in the nucleus on a per cell basis, the proportion of DDX1 protein relative to total protein was similar in both cellular compartments (Fig. 8A). Both the 90- and 84-kDa bands were present in cytosol and nuclear extracts, although the bottom band was more readily apparent in the cytosol. By running the gel for an extended period of time (twice as long as usual), we were able to detect an additional weak band at ~88 kDa in both nuclear and cytosolic extracts.

To determine whether DDX1 consistently localizes to both the cytoplasm and nucleus, we prepared cytosol and nuclear extracts from two additional *DDX1*-amplified lines, Y79 and IMR-32, as well as from nonamplified RB(E)-2, HL60, and HeLa. DDX1 protein was found in both the nucleus and cytoplasm of IMR-32, primarily in the cytoplasm of Y79, and mainly in the nucleus of the three nonamplified lines (Fig. 8B). In addition, DDX1 was almost exclusively found in nuclear

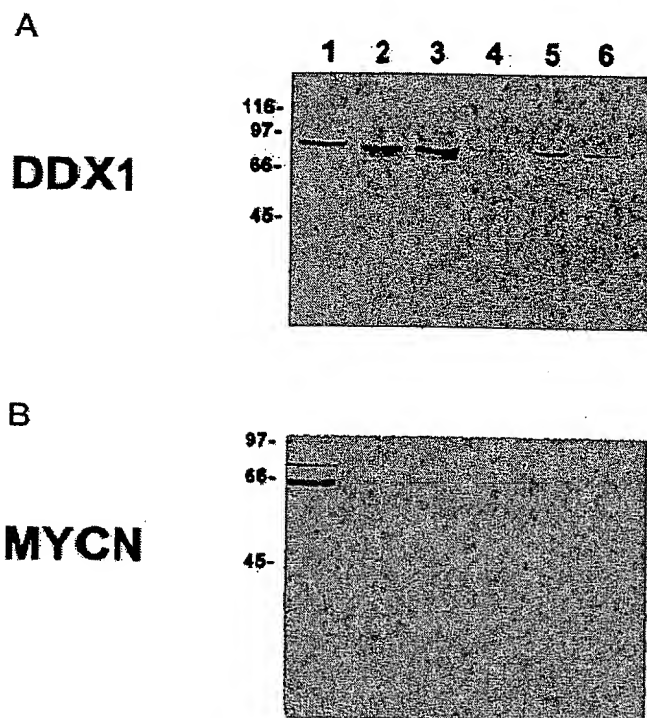


FIG. 9. Subcellular location of DDX1 protein. RB522A cells were fractionated into nuclear (lane 1), S100 and S4 cytosol (lanes 2 and 3), P2 membrane (lane 4), P3 membrane (lane 5), and P4 membrane (lane 6) fractions. Twenty μ g of protein were loaded in each lane and run on a 10% SDS-PAGE gel. A, DDX1 protein was detected using a 1:5000 dilution of the antiserum to the N terminus of DDX1. B, MYCN protein was detected using a commercially available antibody at a 1:200 dilution. Size markers (kilodaltons) are indicated on the side.

extracts prepared from normal GM38 fibroblasts (data not shown). We used anti-actin antibody to ensure that our nuclear and cytosolic extracts were not cross-contaminated (Fig. 8B).

We next carried out a more detailed analysis of DDX1 subcellular location using two different approaches: (i) fractionation of cellular components into nuclei; S100 or S4 cytosol (containing soluble cytoplasmic components, including 40 S ribosomes); P2 (heavy mitochondria, plasma membrane fragments plus material trapped by these membranes); P3 (mitochondria, lysosomes, peroxisomes, Golgi membranes, some rough endoplasmic reticulum); and P4 (microsomes from smooth and rough endoplasmic reticulum, Golgi and plasma membranes) (43); and (ii) immunogold electron microscopy. The DDX1-amplified RB522A cell line was used for both experiments. The fractionation procedures indicate that DDX1 is mainly in the nucleus and in the cytosol (S4 and S100 fractions) of RB522A cells (Fig. 9A). As a control, we used anti-human MYCN antibody to determine the location of MYCN (also amplified in RB522A) in our subcellular fractions. As shown in Fig. 9B, MYCN was primarily found in the nucleus, as one would expect of a transcription factor.

For the electron microscopy analysis, antiserum to the N terminus of DDX1 was coupled to protein A gold particles, and the distribution of DDX1 was examined in RB522A cells fixed in paraformaldehyde and glutaraldehyde. DDX1 was present in both the cytoplasm and nucleus (data not shown). There was no association with either cell organelles or with nuclear or plasma membranes.

DISCUSSION

There are presently few clues as to the function of DDX1 in normal and cancer cells. Our earlier data indicate that DDX1 mRNA is present at higher levels in fetal tissues of neural origin (retina and brain) compared with other fetal tissues (21).

There may therefore be a requirement for elevated levels of this putative RNA helicase for the efficient production or processing of neural specific transcripts. A role in cancer formation or progression is an intriguing possibility, because overexpression of an RNA unwinding protein could affect the secondary structure of RNAs in such a way as to alter the expression of specific proteins in tumor cells. DDX1 is co-amplified with MYCN in a subset of RB and NB cell lines and tumors (37–39). MYCN amplification is common in stage IV NB tumors and is a well documented indicator of poor prognosis. A general trend toward a poorer clinical prognosis is observed when both the MYCN and DDX1 genes are amplified compared with when only MYCN is amplified (38, 39), suggesting a possible role for DDX1 in NB tumor formation or progression.

It is generally accepted that co-amplified genes are not overexpressed unless they provide a selective growth advantage to the cell (48, 49). For example, although ERBA is closely linked to ERBB2 in breast cancer and both genes are commonly amplified in these tumors, ERBA is not overexpressed (48). Similarly, three genes mapping to 12q13–14 (CDK4, SAS, and MDM2) are overexpressed in a high percentage of malignant gliomas showing amplification of this chromosomal region, while other genes mapping to this region (GADD153, GLI, and A2MR) are rarely overexpressed in gene-amplified malignant gliomas (50, 51). The first three genes are probably the main targets of the amplification process, while the latter three genes are probably incidentally included in the amplicons. The data shown here indicate that DDX1 is overexpressed at both the protein and RNA levels in DDX1-amplified RB and NB cell lines and that there is a strong correlation between DDX1 gene copy number, DDX1 RNA levels, and DDX1 protein levels in these lines. Our results are therefore consistent with DDX1 overexpression playing a positive role in some aspect of NB and RB tumor formation or progression. Recently, Weiss *et al.* (52) have shown that transgenic mice that overexpress MYCN develop NB tumors several months after birth. They conclude that MYCN overexpression can contribute to the initiation of tumorigenesis but that additional events are required for tumor formation. Amplification of DDX1 may represent one of many alternative pathways by which a normal precursor “neuroblast” or “retinoblast” cell gains malignant properties.

The function of the majority of tissue-specific or developmentally regulated DEAD box genes remains unknown. However, some members of this protein family have been either directly or indirectly implicated in tumorigenesis. For example, the p68 gene has been found to be mutated in the ultraviolet light-induced murine tumor 8101 (53), while DDX6 (also known as RCK or p54) is encoded by a gene located at the breakpoint of the translocation involving chromosomes 11 and 14 in a cell line derived from a B-cell lymphoma (54, 55). Similarly, the production of a chimeric protein between DDX10 and the nucleoporin gene NUP98 has been proposed to be involved in the pathogenesis of a subset of myeloid malignancies with inv(11) (p15q22) (56). Interestingly, Grandori *et al.* (57) have shown that MYCC interacts with a DEAD box gene called MrDb, suggesting that the transcription of some DEAD box genes could be regulated through interaction with members of the MYC family. Future work will involve determining whether DDX1 represents another member of the DEAD box family with a role in the tumorigenic process.

DEAD box proteins have been implicated in translation initiation, RNA splicing, RNA degradation, and RNA stability (3, 18, 19). We carried out subcellular localization studies in an attempt to obtain a general indication of the function of DDX1. We found DDX1 protein in both the cytoplasm and nucleus of DDX1-amplified NB and RB lines. In contrast, DDX1 was

mainly located in the nucleus of nonamplified cell lines and normal fibroblast cultures. DDX1 was not associated with cellular organelles or with membranes based on immunoelectron microscopy. We therefore propose that the primary role of DDX1 is in the nucleus. The presence of DDX1 in the cytoplasm of DDX1-amplified cells may indicate that the amount of DDX1 protein that is allowed in the nucleus is tightly regulated. Alternatively, DDX1 may play a dual role in the nucleus and cytoplasm of DDX1-amplified cells.

An important component of our analysis was to identify the translation and transcription initiation sites of DDX1. We used a combination of techniques to identify the transcription start site: screening of RB and fetal brain libraries, RACE, primer extension, genomic DNA sequencing, S1 nuclease mapping, and Northern blot analysis using probes to the predicted 5'-end of the transcript. The transcription start site identified using these techniques is located ~300 nt upstream of the predicted translation initiation codon and was readily detected in three DDX1-amplified lines and barely detectable in a fourth amplified line. The 5'-untranslated region as well as the first in frame methionine are encoded within the first exon of DDX1. An in frame stop codon is located 123 nt upstream of the predicted initiation codon. We were unable to identify the transcription initiation site of DDX1 in two of the six amplified lines tested as well as in nonamplified lines. Although it remains possible that there are different transcription start sites in different cell lines, detection of lower-levels (rather than the absence) of the 5'-most 160 nt of the DDX1 transcript in IMR-32, Y79, and LA-N-5 compared with RB522A, BE(2)-C, and LA-N-1 supports a quantitative rather than a qualitative difference in the 5'-end of this transcript in these cells. Our results suggest that the 5'-end of DDX1 mRNA is rarely intact, even in mRNA preparations that otherwise appear to be of high quality based on analysis of control transcripts. The 5'-end of DDX1 mRNA may therefore be especially susceptible to degradation, perhaps because of its sequence and/or secondary structure.

In conclusion, we have mapped the 5'-end of the 2.7-kb DDX1 transcript and have identified the predicted translation initiation site of DDX1 protein. We have found that DDX1-amplified RB and NB tumor lines overexpress DDX1 protein and that there is a good correlation between gene copy number and both transcript and protein levels in these cells. We have shown that DDX1 protein is primarily located in the nucleus of cells that are not DDX1-amplified. In contrast, DDX1 is present in both the nucleus and cytoplasm of DDX1-amplified NB and RB lines. A cytoplasmic location in DDX1-amplified lines may indicate that the amount of nuclear DDX1 is tightly regulated or that DDX1 plays a dual role in the cytoplasm and nucleus of these cells.

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Aneuploidy and cancer

Subrata Sen, PhD

Numeric aberrations in chromosomes, referred to as aneuploidy, is commonly observed in human cancer. Whether aneuploidy is a cause or consequence of cancer has long been debated. Three lines of evidence now make a compelling case for aneuploidy being a discrete chromosome mutation event that contributes to malignant transformation and progression process. First, precise assay of chromosome aneuploidy in several primary tumors with *in situ* hybridization and comparative genomic hybridization techniques have revealed that specific chromosome aneusomies correlate with distinct tumor phenotypes. Second, aneuploid tumor cell lines and *in vitro* transformed rodent cells have been reported to display an elevated rate of chromosome instability, thereby indicating that aneuploidy is a dynamic chromosome mutation event associated with transformation of cells. Third, and most important, a number of mitotic genes regulating chromosome segregation have been found mutated in human cancer cells, implicating such mutations in induction of aneuploidy in tumors. Some of these gene mutations, possibly allowing unequal segregations of chromosomes, also cause tumorigenic transformation of cells *in vitro*. In this review, the recent publications investigating aneuploidy in human cancers, rate of chromosome instability in aneuploidy tumor cells, and genes implicated in regulating chromosome segregation found mutated in cancer cells are discussed. *Curr Opin Oncol* 2000, 12:82-88 © 2000 Lippincott Williams & Wilkins, Inc.

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Abbreviations

CGH . comparative genomic hybridization
CHE Chinese hamster embryo cells
FISH fluorescence *in situ* hybridization
HPRC hereditary papillary renal carcinoma
ISH *in situ* hybridization

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Cancer research over the past decade has firmly established that malignant cells accumulate a large number of genetic mutations that affect differentiation, proliferation, and cell death processes. In addition, it is also recognized that most cancers are clonal, although they display extensive heterogeneity with respect to karyotypes and phenotypes of individual clonal populations. It is estimated that numeric chromosomal imbalance, referred to as *aneuploidy*, is the most prevalent genetic change recorded among over 20,000 solid tumors analyzed thus far [1]. Phenotypic diversity of the clonal populations in individual tumors involve differences in morphology, proliferative properties, antigen expression, drug sensitivity, and metastatic potentials. It has been proposed that an underlying acquired genetic instability is responsible for the multiple mutations detected in cancer cells that lead to tumor heterogeneity and progression [2]. In a somewhat contradictory argument, it has also been suggested that clonal expansion due to selection of cells undergoing normal rates of mutation can explain malignant transformation and progression process in humans [3]. Acquired genetic instability, nonetheless, is considered important for more rapid progression of the disease [4•]. Although the original hypothesis on genetic instability in cancer primarily focused on chromosome imbalances in the form of aneuploidy in tumor cells, the actual relevance of such mutations in cancer remains a controversial issue.

Whether or not aneuploidy contributes to the malignant transformation and progression process has long been debated. A prevalent idea on genetics of cancer referred to as "somatic gene mutation hypothesis" contends that gene mutations at the nucleotide level alone can cause cancer by either activating cellular proto-oncogenes to dominant cancer causing oncogenes and/or by inactivating growth inhibitory tumor suppressor genes. In this scheme of things chromosomal instability in the form of aneuploidy is a mere consequence rather than a cause of malignant transformation and progression process.

In this review, some of the recent observations on the subject are discussed and compelling evidence is provided to suggest that aneuploidy is a distinct form of genetic instability in cancer that frequently correlates with specific phenotypes and stages of the disease. Furthermore, discrete genetic targets affecting chromosomal stability in cancer cells, recently identified, are also discussed. These data provide a new direction toward elucidating the molecular mechanisms responsi-

ble for induction of aneuploidy in cancer and may eventually be exploited as novel therapeutic targets in the future.

Genetic alterations in cancer

Alterations in many genetic loci regulating growth, senescence, and apoptosis, identified in tumor cells, have led to the current understanding of cancer as a genetic disease. The genetic changes identified in tumors include: subtle mutations in genes at the nucleotide level; chromosomal translocations leading to structural rearrangements in genes; and numeric changes in either partial segments of chromosomes or whole chromosomes (aneuploidy) causing imbalance in gene dosage.

For the purpose of this review, both segmental and whole chromosome imbalances leading to altered DNA dosage in cancer cells are included as examples of aneuploidy.

Incidence of aneuploidy in cancer

Evidence of aneuploidy involving one or more chromosomes have been commonly reported in human tumors. Although these observations were initially made using classic cytogenetic techniques late in a tumor's evolution and were difficult to correlate with cancer progression, more recent studies have reported association of specific nonrandom chromosome aneuploidy with different biologic properties such as loss of hormone dependence and metastatic potential [5].

Classic cytogenetic studies performed on tumor cells had serious limitations in scope because they were applicable only to those cases in which mitotic chromosomes could be obtained. Because of low spontaneous rates of cell division in primary tumors, analyses depended on cells either derived selectively from advanced metastases or those grown *in vitro* for variable periods of time. In both instances, metaphases analyzed represented only a subset of primary tumor cell population. Two major advances in cytogenetic analytic techniques, *in situ* hybridization (ISH) and comparative genomic hybridization (CGH), have allowed better resolution of chromosomal aberrations in freshly isolated tumor cells [6]. ISH analyses with chromosome-specific DNA probes, a powerful adjunct to metaphasic analysis, allows assessment of chromosomal anomalies within tumor cell populations in the contexts of whole nuclear architecture and tissue organization. CGH allows genome wide screening of chromosomal anomalies without the use of specific probes even in the absence of prior knowledge of chromosomes involved. Although both techniques have certain limitations in terms of their resolution power, they nonetheless provide a better approximation of chromosomal changes occurring among tumors of various histology, grade, and stage

compared with what was possible with the classic cytogenetic techniques. Genomic ploidy measurements have also been performed at the DNA level with flow cytometry and cytofluorometric methods. Although these assays underestimate chromosome ploidy due to a chromosomal gain occasionally masking a chromosomal loss in the same cell, several studies using these methods have supported the conclusion that DNA aneuploidy closely associates with poor prognosis in various cancers [7,8]. This discussion of some recent examples published on aneuploidy in cancer includes discussion of studies dealing with DNA ploidy measurements as well. Most of these observations are correlative without direct proof of specific involvement of genes on the respective chromosomes. Identification of putative oncogenes and tumor suppressor genes on gained and lost chromosomes in aneuploid tumors, however, are providing strong evidence that chromosomes involved in aneuploidy play a critical role in the tumorigenic process.

In renal tumors, either segmental or whole chromosome aneuploidy appears to be uniquely associated with specific histologic subtypes [9]. Tumors from patients with hereditary papillary renal carcinomas (HPRC) commonly show trisomy of chromosome 7, when analyzed by CGH. Germline mutations of a putative oncogene *MET* have been detected in patients with HPRC. A recent study [10] has demonstrated that an extra copy of chromosome 7 results in nonrandom duplication of the mutant *MET* allele in HPRC, thereby implicating this trisomy in tumorigenesis. The study suggested that mutation of *MET* may render the cells more susceptible to errors in chromosome replication, and that clonal expansion of cells harboring duplicated chromosome 7 reflects their proliferative advantage. In addition to chromosome 7, trisomy of chromosome 17 in papillary tumors and also of chromosome 8 in mesoblastic nephroma are commonly seen. Association of specific chromosome imbalances with benign and malignant forms of papillary renal tumors, therefore, not only contribute to an understanding of tumor origins and evolution, but also implicate aneuploidy of the respective chromosomes in the tumorigenic transformation process.

In colorectal tumors, chromosome aneuploidy is a common occurrence. In fact, molecular allelotyping studies have suggested that limited karyotyping data available from these tumors actually underestimate the true extent of these changes. Losses of heterozygosity reflecting loss of the maternal or paternal allele in tumors are widespread and often accompanied by a gain of the opposite allele. Therefore, for example, a tumor could lose a maternal chromosome while duplicating the same paternal chromosome, leaving the tumor cell

with a normal karyotype and ploidy but an aberrant allelotype. It has been estimated that cancer of the colon, breast, pancreas, or prostate may lose an average of 25% of its alleles. It is not unusual to discover that a tumor has lost over half of its alleles [4]. In clinical settings, DNA ploidy measurements have revealed that DNA aneuploidy indicates high risk of developing severe premalignant changes in patients with ulcerative colitis, who are known to have an increased risk of developing colorectal cancer [11]. DNA aneuploidy has been found to be one of the useful indicators of lymph node metastasis in patients with gastric carcinoma and associated with poor outcome compared with diploid cases [12,13]. CGH analyses of chromosome aneuploidy, on the other hand, was reported to correlate gain of chromosome 20q with high tumor S phase fractions and loss of 4q with low tumor apoptotic indices [14]. Aneuploidy of chromosome 4 in metastatic colorectal cancer has recently been confirmed in studies that used unbiased DNA fingerprinting with arbitrarily primed polymerase chain reactions to detect moderate gains and losses of specific chromosomal DNA sequences [15]. The molecular karyotype (amplotype) generated from colorectal cancer revealed that moderate gains of sequences from chromosomes 8 and 13 occurred in most tumors, suggesting that overrepresentation of these chromosomal regions is a critical step for metastatic colorectal cancer.

In addition to being implicated in tumorigenesis and correlated with distinct tumor phenotypes, chromosome aneuploidy has been used as a marker of risk assessment and prognosis in several other cancers. The potential value of aneuploidy as a noninvasive tool to identify individuals at high risk of developing head and neck cancer appears especially promising. Interphase fluorescence *in situ* hybridization (FISH) revealed extensive aneuploidy in tumors from patients with head and neck squamous cell carcinomas (HNSCC) and also in clinically normal distant oral regions from the same individuals [16,17]. It has been proposed that a panel of chromosome probes in FISH analyses may serve as an important tool to detect subclinical tumorigenesis and for diagnosis of residual disease. The presence of aneuploid or tetraploid populations is seen in 90% to 95% of esophageal adenocarcinomas, and when seen in conjunction with Barrett's esophagus, a premalignant condition, predicts progression of disease [18,19]. Chromosome ploidy analyses in conjunction with loss of heterozygosity and gene mutation studies in Barrett's esophagus reflect evolution of neoplastic cell lineages *in vivo* [20]. Evolution of neoplastic progeny from Barrett's esophagus following somatic genetic mutations frequently involves bifurcations and loss of heterozygosity at several chromosomal loci leading to aneuploidy and cancer. Accordingly, it is hypothesized that during

tumor cell evolution diploid cell progenitors with somatic genetic abnormalities undergo expansion with acquired genetic instability. Such instability, often manifested in the form of increased incidence of aneuploidy, enters a phase of clonal evolution beginning in premalignant cells that proceeds over a period of time and occasionally leads to malignant transformation. The clonal evolution continues even after the emergence of cancer.

The significance of DNA and chromosome aneuploidy in other human cancers continue to be evaluated. Among papillary thyroid carcinomas, aneuploid DNA content in tumor cells was reported to correlate with distant metastases, reflecting worsened prognosis [21]. Genome wide screening of follicular thyroid tumors by CGH, on the other hand, revealed frequent loss of chromosome 22 in widely invasive follicular carcinomas [22]. Chromosome copy number gains in invasive neoplasm compared with foci of ductal carcinoma *in situ* (DCIS) with similar histology have been proposed to indicate involvement of aneuploidy in progression of human breast cancer [23]. ISH analyses of cervical intraepithelial neoplasia has provided suggestive evidence that chromosomes 1, 7 and X aneusomy is associated with progression toward cervical carcinoma [24].

Although the prognostic value of numeric aberrations remains a matter of debate in human hematopoietic neoplasia, there have been recent studies to suggest that the presence of monosomy 7 defines a distinct subgroup of acute myeloid leukemia patients [25]. It is interesting in this context that therapy-related myelodysplastic syndromes have been reported to display monosomy 5 and 7 karyotypes, reflecting poor prognosis [26].

The clinical observations, mentioned previously, are supported by *in vitro* studies in human and rodent cells in which aneuploidy is induced at early stages of transformation [27,28]. It is even suggested that aneuploidy may cause cell immortalization, in some instances, that is a critical step preceding transformation.

Finally, in an interesting study to develop transgenic mouse models of human chromosomal diseases, chromosome segment specific duplication and deletions of the genome were reported to be constructed in mouse embryonic stem cells [29]. Three duplications for a portion of mouse chromosome 11 syntenic with human chromosome 17 were established in the mouse germline. Mice with 1Mb duplication developed corneal hyperplasia and thymic tumors. The findings represent the first transgenic mouse model of aneuploidy of a defined chromosome segment that documents the direct role of chromosome aneusomy in tumorigenesis.

Aneuploidy as "dynamic cancer-causing mutation" instead of a "consequential state" in cancer

According to the hypothesis previously discussed, aneuploidy represents either a "gain of function" or "loss of function" mutation at the chromosome level with a causative influence on the tumorigenesis process. The hypothesis, however, is based only on circumstantial evidence even though existence of aneuploidy is correlated with different tumor phenotypes. The existence of numeric chromosomal alterations in a tumor does not mean that the change arose as a dynamic mutation due to genomic instability, because several factors could lead to consequential aneuploidy in tumors, also. Although aneuploidy as a dynamic mutation due to genomic instability in tumor cells would occur at a certain measurable rate per cell generation, a consequential state of aneuploidy in tumors may not occur at a predictable rate under similar conditions or in tumors with similar phenotypes. In addition to genomic instability, differences in environmental factors with selective pressure, could explain high incidence of aneuploidy and other somatic mutations in tumors compared with normal cells [4]. These include humoral, cell substratum, and cell-cell interaction differences between tumor and normal cell environments. It could be argued that despite similar rates of spontaneous aneuploidy induction in normal and tumor cells, the latter are selected to proliferate due to altered selective pressure in the tumor cell environment, whereas the normal cells are eliminated through activation of apoptosis. Alternatively, of course, one could postulate that selective expression or overexpression of anti-apoptotic proteins or inactivation of proapoptotic proteins in tumor cells may counteract default induction of apoptosis in G2/M phase cells undergoing missegregation of chromosomes. Recent demonstration of overexpression of a G2/M phase anti-apoptotic protein survivin in cancer cells [30] suggests that this protein may favor aberrant progression of aneuploid transformed cells through mitosis. This would then lead to proliferation of aneuploid cell lineages, which may undergo clonal evolution.

To ascertain that aneuploidy is a dynamic mutational event, various human tumor cell lines and transformed rodent cell lines have been analyzed for the rate of aneuploidy induction. When grown under controlled *in vitro* conditions, such conditions ensure that environmental factors do not influence selective proliferation of cells with chromosome instability. In one study, Lengauer *et al.* [31•] provided unequivocal evidence by FISH analyses that losses or gains of multiple chromosomes occurred in excess of 10^{-2} per chromosome per generation in aneuploid colorectal cancer cell lines. The study further concluded that such chromosomal instability appeared to be a dominant trait. Using another *in*

vitro model system of Chinese hamster embryo (CHE) cells, Duesberg *et al.* [32•] have also obtained similar results. With clonal cultures of CHE cells, transformed with nongenotoxic chemicals and a mitotic inhibitor, these authors demonstrated that the overwhelming majority of the transformed colonies contained more than 50% aneuploid cells, indicating that aneuploidy would have originated from the same cells that underwent transformation. All the transformed colonies tested were tumorigenic. It was further documented that the ploidy factor representing the quotient of the modal chromosome number divided by the normal diploid number, in each clone, correlated directly with the degree of chromosomal instability. Therefore, chromosomal instability was found proportional to the degree of aneuploidy in the transformed cells and the authors hypothesized that aneuploidy is a unique mechanism of simultaneously altering and destabilizing, in a massive manner, the normal cellular phenotypes. In the absence of any evidence that the transforming chemicals used in the study did not induce other somatic mutations, it is difficult to rule out the contribution of such mutations in the transformation process. These results nonetheless make a strong case for aneuploidy being a dynamic chromosome mutation event intimately associated with cancer.

Aneuploidy versus somatic gene mutation in cancer

The idea that numeric chromosome imbalance or aneuploidy is a direct cause of cancer was proposed at the turn of the century by Theodore Boveri [33]. However, the hypothesis was largely ignored over the last several decades in favor of the somatic gene mutation hypothesis, mentioned earlier. Evidence accumulating in the literature lately on specific chromosome aneusomies recognized in primary tumors, incidence of aneuploidy in cells undergoing transformation, and aneuploid tumor cells showing a high rate of chromosome instability have led to the rejuvenation of Boveri's hypothesis. The concept has recently been discussed as a "vintage wine in a new bottle" [34•]. The author points out that except for rare cancers caused by dominant retroviral oncogenes, diploidy does not seem to occur in solid tumors, whereas aneuploidy is a rule rather than exception in cancer.

Aneuploidy as an effective mutagenic mechanism driving tumor progression, on the other hand, is being recognized as a viable solution to the paradox that with known mutation rate in non-germline cells ($\sim 10^{-7}$ per gene per cell generation) tumor cell lineages cannot accumulate enough mutant genes during a human lifetime [35]. The concept is gaining significant credibility since genes that potentially affect chromosome segregation were found mutated in human cancer. Some of

these genes have also been shown to have transforming capability in *in vitro* assays. Selected recent publications describing the findings are being discussed below in reference to the mitotic targets potentially involved in inducing chromosome segregation anomalies in cells.

Potential mitotic targets and molecular mechanisms of aneuploidy

Because aneuploidy represents numeric imbalance in chromosomes, it is reasonable to expect that aneuploidy arises due to missegregation of chromosomes during cell division. There are many potential mitotic targets, which could cause unequal segregation of chromosomes (Fig. 1). Recent investigations have identified several genes involved in regulating these mitotic targets and mitotic checkpoint functions, which can be implicated in induction of aneuploidy in tumor cells. This discussion is restricted to those mitotic targets and checkpoint genes whose abnormal functioning has been observed in cancer or has been shown to cause tumorigenic transformation of cells, in recent years. The role of telomeres is discussed elsewhere in this issue. For a more detailed description of the components of mitotic machinery and their possible involvement in causing chromosome segregation abnormalities in tumor cells, readers may refer to a recently published review [36•].

Among the mitotic targets implicated in cancer, centrosome defects have been observed in a wide variety of malignant human tumors. Centrosomes play a central role in organizing the microtubule network in interphase cells and mitotic spindle during cell division. Multipolar mitotic spindles have been observed in human cancers *in situ* and abnormalities in the form of supernumerary

centrosomes, centrosomes of aberrant size and shape as well as aberrant phosphorylation of centrosome proteins have been reported in prostate, colon, brain, and breast tumors [37,38]. In view of the findings that abnormal centrosomes retain the ability to nucleate microtubules *in vitro*, it is conceivable that cells with abnormal centrosomes may missegregate chromosomes producing aneuploid cells. The molecular and genetic bases of abnormal centrosome generation and the precise pathway through which they regulate the chromosome segregation process remain to be elucidated. Recent discovery of a centrosome-associated kinase STK15/BTAK/aurora2, naturally amplified and overexpressed in human cancers, has raised the interesting possibility that aberrant expression of this kinase is critically involved in abnormal centrosome function and unequal chromosome segregation in tumor cells [39,40]. Exogenous expression of the kinase in rodent and human cells was found to correlate with an abnormal number of centrosomes, unequal partitioning of chromosomes during division, and tumorigenic transformation of cells. It is relevant in this context to mention that the *Xenopus* homologue of human STK15/BTAK/aurora2 kinase has recently been shown to phosphorylate a microtubule motor protein XIEg5, the human orthologue of which is known to participate in the centrosome separation during mitosis [41]. Findings on STK15/aurora2 kinase, thus, provide an interesting lead to a possible molecular mechanism of centrosome's role in oncogenesis. Centrosomes have, of late, been implicated in oncogenesis from studies revealing supernumerary centrosomes in β 53-deficient fibroblasts and overexpression of another centrosome kinase PLK1 being detected in human non-small cell lung cancer [42].

One of the critical events that ensures equal partitioning of the chromosomes during mitosis is the proper and timely separation of sister chromatids that are attached to each other and to the mitotic spindle. Untimely separation of sister chromatids has been suspected as a cause of aneuploidy in human tumors. Cohesion between sister chromatids is established during replication of chromosomes and is retained until the next metaphase/anaphase transition. It has been shown that during metaphase-anaphase transition, the anaphase promoting complex/cyclosome triggers the degradation of a group of proteins called securins that inhibit sister chromatid separation. A vertebrate securin (v-securin) has recently been identified that inhibits sister chromatid separation and is involved in transformation and tumorigenesis. Subsequent analysis revealed that the human securin is identical to the product of the gene called pituitary tumor transforming gene, which is overexpressed in some tumors and exhibits transforming activity in NIH3T3 cells. It is proposed that elevated expression of the v-securin may contribute to generation of malignant tumors due to

Figure 1. Potential mitotic targets causing aneuploidy in oncogenesis

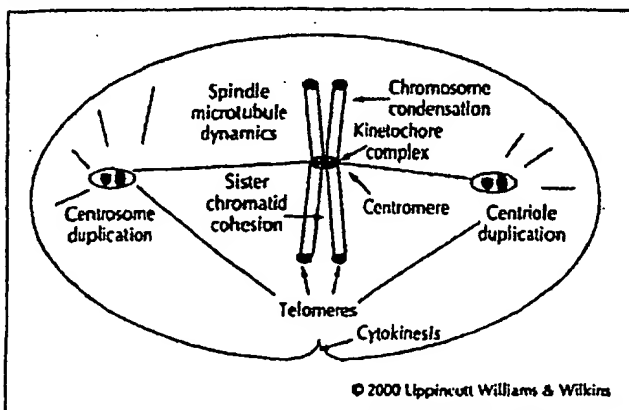


Diagram illustrates that defects in several processes involving chromosomal, spindle microtubule, and centrosomal targets, in addition to abnormal cytokinesis, may cause unequal partitioning of chromosomes during mitosis, leading to aneuploidy. Recently obtained evidence in favor of some of these possibilities is discussed in the text.

chromosome gain or loss produced by errors in chromatid separation [43•].

Normal progression through mitosis during prophase to anaphase transition is monitored at least at two checkpoints: One checkpoint operates during early prophase at G2 to metaphase progression while the second ensures proper segregation of chromosomes during metaphase to anaphase transition. Several mitotic checkpoint genes responding to mitotic spindle defects have been identified in yeast. The metaphase-anaphase transition is delayed following activation of this checkpoint during which kinetochores remain unattached to the spindle. The signal is transmitted through a kinetochore protein complex consisting of Mps1p and several Mad and Bub proteins [44]. It is expected that for unequal chromosome segregation to be perpetuated through cell proliferation cycles giving rise to aneuploidy, checkpoint controls have to be abrogated.

Following this logic, Vogelstein *et al.* [45•] hypothesized that aneuploid tumors would reveal mutation in mitotic spindle checkpoint genes. Subsequent studies by these investigators have proven the validity of this hypothesis and a small fraction of human colorectal cancers have revealed the presence of mutations in either hBub1 or hBubR1 checkpoint genes. It was further revealed that mutant hBUB1 could function in a dominant negative manner conferring an abnormal spindle checkpoint when expressed exogenously. Inactivation of spindle checkpoint function in virally induced leukemia has also recently been documented following the finding that hMAD1 checkpoint protein is targeted by the Tax protein of the human T-cell leukemia virus type 1. Abrogation of hMAD1 function leads to multinucleation and aneuploidy [46].

In addition to mitotic spindle checkpoint defects, failed DNA damage checkpoint function in yeast is frequently associated with aberrant chromosome segregation as well. It, therefore, appears intriguing yet relevant that the human *BRCA1* gene, proposed to be involved in DNA damage checkpoint function, when mutated by a targeted deletion of exon 11 led to defective G2/M cell cycle checkpoint function and genetic instability in mouse embryonic fibroblasts [47]. The cells revealed multiple functional centrosomes and unequal chromosome segregation and aneuploidy. Although the molecular basis for these abnormalities is not known at this time, it raises the interesting possibility that such an aneuploidy-driven mechanism may be involved in tumorigenesis in individuals carrying germline mutations of *BRCA1* gene.

Conclusion

Growing evidence from human tumor cytogenetic investigations strongly suggest that aneuploidy is associated with the development of tumor phenotypes. Clinical findings of correlation between aneuploidy and tumorigenesis are supported by studies with *in vitro* grown transformed cell lines. Molecular genetic analyses of tumor cells provide credible evidence that mutations in genes controlling chromosome segregation during mitosis play a critical role in causing chromosome instability leading to aneuploidy in cancer. Further elucidation of molecular and physiologic bases of chromosome instability and aneuploidy induction could lead to the development of new therapeutic approaches for common forms of cancer.

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ORIGINAL ARTICLE

Identification of putative oncogenes in lung adenocarcinoma by a comprehensive functional genomic approach

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Amplification and overexpression of putative oncogenes confer growth advantages for tumor development. We used a functional genomic approach that integrated simultaneous genomic and transcript microarray, proteomics, and tissue microarray analyses to directly identify putative oncogenes in lung adenocarcinoma. We first identified 183 genes with increases in both genomic copy number and transcript in six lung adenocarcinoma cell lines. Next, we used two-dimensional polyacrylamide gel electrophoresis and mass spectrometry to identify 42 proteins that were overexpressed in the cancer cells relative to normal cells. Comparing the 183 genes with the 42 proteins, we identified four genes – *PRDX1*, *EEF1A2*, *CALR*, and *KCIP-1* – in which elevated protein expression correlated with both increased DNA copy number and increased transcript levels (all $r > 0.84$, two-sided $P < 0.05$). These findings were validated by Southern, Northern, and Western blotting. Specific inhibition of *EEF1A2* and *KCIP-1* expression with siRNA in the four cell lines tested suppressed proliferation and induced apoptosis. Parallel fluorescence *in situ* hybridization and immunohistochemical analyses of *EEF1A2* and *KCIP-1* in tissue microarrays from patients with lung adenocarcinoma showed that gene amplification was associated with high protein expression for both genes and that protein overexpression was related to tumor grade, disease stage, Ki-67 expression, and a shorter survival of patients. The amplification of *EEF1A2* and *KCIP-1* and the presence of overexpressed protein in tumor samples strongly suggest that these genes could be oncogenes and hence potential targets for diagnosis and therapy in lung adenocarcinoma. *Oncogene* (2006) 25, 2628–2635. doi:10.1038/sj.onc.1209289; published online 12 December 2005

Keywords: lung cancer; microarrays; proteomics; tissue microarray

Introduction

In lung adenocarcinoma, as in other types of cancer, gene amplification and the consequent overexpression of the amplified oncogene play an important role in the development of tumors, because their overexpression confers a growth advantage. The ability to identify putative oncogenes that are activated during tumorigenesis could facilitate the choice of molecular genetic targets for diagnosis and therapy of the disease. This concept has been exemplified by *HER-2*, which was first found to be amplified in neuroblastomas and subsequently shown to be associated with poor prognosis in breast cancer (Ross and Fletcher, 1999). Now, *HER-2* aberrations are used as a predictor of response to therapy, and treatment of *HER-2*-positive breast cancer with the monoclonal anti-*HER-2* antibody trastuzumab has been shown to improve prognosis (Ross and Fletcher, 1999). Emerging evidence of common amplicons in lung adenocarcinomas (Luk *et al.*, 2001; Jiang *et al.*, 2004; Tonon *et al.*, 2005) suggests that additional oncogenes remain to be identified; however, conventional techniques are ineffective in pinpointing such oncogenes. Parallel measurement of DNA copy number and mRNA levels in cDNA microarrays permits changes in copy number to be compared with transcription levels on a gene-by-gene basis to generate lists of candidate genes within the defining amplicons (Hyman *et al.*, 2002; Pollack *et al.*, 2002). However, use of transcript patterns does not allow assessment of the expression of protein products or identification of proto-oncogenes. Another approach, identifying differentially expressed proteins by proteomic analysis and then comparing the proteins present with mRNA expression in cDNA microarrays from the same specimens, can clarify the extent to which changes in transcript patterns reflect changes in their cognate proteins and post-transcriptional mechanisms (Chen *et al.*, 2002), but this approach cannot be used to identify oncogenes driven by extensive increases of their gene copy number. Moreover, using individual microarrays or proteomic approaches alone cannot distinguish the cancer-driving oncogenes that directly propel tumor progression from the larger number of passenger genes that may be concurrently over-represented but are not biologically relevant in tumor development.

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In this study, we used a comprehensive approach that integrated simultaneous comparative genomic hybridization (CGH) and transcript microarray with proteomic analyses of six lung adenocarcinoma cell lines. We directly and specifically identified four putative oncogenes that could have been activated through amplification and consequent elevation of transcript expression. We used small interfering RNA (siRNA) to inhibit the expression of two of these four genes in the lung cancer cell lines, which further implicated them in oncogenesis. We then explored the clinical significance of these findings by assessing the expression of these two genes in tissue microarrays of human lung cancer specimens. Our findings underscore the power of integrated functional genomic analyses for identifying putative oncogenes in tumorigenesis; such activated genes could be useful as targets for diagnosis or therapy in lung cancer.

Results

Simultaneous global genomic and transcript analyses identify 183 genes with increases in genomic copy numbers and transcript expression levels

To identify genes in which increased DNA copy number might contribute to increased transcript in lung adenocarcinomas, first we used CGH with microarrays of six lung adenocarcinoma cell lines. We identified 587 genes showing increases in DNA copy number across all six cell lines (Supplementary Table 1S), which were distributed as 90 amplicons on all chromosomes except for chromosomes 13 and Y (Supplementary Table 2S). A subsequent transcript test with the identical arrays of the same cell lines revealed 275 genes that showed increased mRNA levels (Supplementary Table 3S). Using random permutation tests across all cancer cell lines, we identified 183 genes (31%) that showed elevated transcript levels from the 587 genes that were over-represented in the genome (Table 1), suggesting that elevated transcript levels of the 183 genes may reflect their genomic over-representation in the cancer cells. These findings are consistent with previous reports linking genomic changes with altered transcript patterns in breast cancer (Hyman *et al.*, 2002; Pollack *et al.*, 2002). However, our finding that only 31% of the genes showing increased DNA copy numbers had cognate increases in transcript expression in lung adenocarcinomas is different from the overall rates of 40–60% reported for breast cancer (Hyman *et al.*, 2002; Pollack *et al.*, 2002). This discordance may reflect methodologic differences between studies or biological differences between breast cancer and lung adenocarcinoma.

Proteomic analyses identify four genes for which protein abundance was associated with increases in the cognate gene and transcript levels

Analysis of transcript patterns is insufficient for understanding the expression of protein products and the effect of genomic over-representation on the expression

of their cognate proteins. To extend these findings beyond genomic over-representation to expression of the protein products of those genes, we next assessed protein expression in the same cell lines by two-dimensional polyacrylamide gel electrophoresis (PAGE) and found that 42 different proteins, representing 42 individual genes, were significantly increased in the cancer cell lines (Table 2; Supplementary Figures 1S and 2S). Some of these proteins were identified as having multiple isoforms, and all individual isoforms exhibited increases in expression ranging from 4.6 to 12.8 times their expression in normal lung tissue cells. In comparing protein level of the 42 genes with changes in their cognate genomic and mRNA expression from the global microarray analyses, we found that four (9.5%) of those 42 genes – *PRDX1*, *EEF1A2*, *CALR*, and *KCIP-1* – showed statistically significant correlations between elevated protein expression and increases in both copy number and mRNA expression (all $r > 0.84$; $P < 0.05$) (Table 2) in the cancer cell lines. These findings imply that the abundance of these four proteins is attributable to the amplification and consequent elevated transcription of their cognate genes.

Validation of copy number, transcript, and protein expression of PRDX1, EEF1A2, CALR, and KCIP-1 in lung cancer cell lines

To confirm our findings from the high-throughput analyses, we next used Southern, Northern, and Western blotting to assess DNA, RNA, and protein levels for the four genes identified in the six cell lines. For comparison, we arbitrarily chose one gene, *NFKB1*, in which an increase in protein level did not correlate with genetic changes. Overall, we found excellent concordance between the CGH microarray and Southern blotting analyses, transcript array and Northern blotting analyses, and proteomic and Western blotting analyses for all five genes (Figure 1). For example, *KCIP-1* showed fivefold amplification in five of the six cancer cell lines, whereas *NFKB1* showed no such increase in any of the cell lines. As for transcript expression, Northern blotting of *EEF1A2* showed high expression in five of the six cancer cell lines; again, levels of *NFKB1* transcript were not increased in any cancer cell line as compared with normal bronchial epithelial cells. The results of Western blotting were also consistent with the results of the proteomic experiments; for example, five of the cancer cell lines exhibited strong protein bands for *PRDX1* as compared with normal cells. These findings provide strong support for the validity of the results derived from the high-throughput techniques in this study.

These parallel analyses also revealed close correlations in the extent of changes in gene copies, transcript, and protein of each of the four genes in the cancer cell lines. For example, in the five cancer cell lines that showed at least fourfold increases in *EEF1A2* copy number, expression of transcript and protein was also increased by at least a factor of four as well (relative to their expression in normal cells) (Supplementary Figure 3S). The protein abundance of the four genes showing

Table 1 List of 183 genes with statistically significant correlation (0.05) between genomic copy number and transcript level

Gene symbol	Chro.	Distance from p arm of each chromosome (Mb)	α
ENO1	1	8.5	0.0085
DDOST	1	20.1	0.0111
SFN	1	26.4	0.0113
MLP	1	32.2	0.0114
AKR1A1	1	45.4	0.0128
PRDX1	1	45.4	0.0122
UQCRH	1	46.2	0.0125
RPL7	1	96.4	0.0127
COL11A1	1	102.6	0.0129
MCL1	1	147.3	0.0222
PSMB4	1	148.1	0.0131
JTB	1	150.7	0.0134
RPS27	1	150.7	0.0135
HAX1	1	151	0.0266
MUC1	1	151.9	0.0143
CCT3	1	153.1	0.0167
CRABP2	1	153.4	0.0148
TKT	1	159.3	0.0152
ATP1B1	1	165.8	0.0234
CHIT1	1	199.7	0.0154
SNRPE	1	200.2	0.0165
YWHAQ	2	9.6	0.0159
ODCI	2	10.60	0.0119
RPL31	2	101.20	0.0161
BENE	2	110.40	0.0169
STAT1	2	191.80	0.0175
HSPD1	2	198.30	0.0277
HSPE1	2	198.30	0.0185
RPL37A	2	217.30	0.0388
IGFBP2	2	217.50	0.0189
RPS7	2	3.30	0.0193
RAB1A	2	65.30	0.0204
IGKC	2	89.00	0.0285
LTF	3	46.3	0.0455
PFN2	3	151	0.0207
KPNA4	3	161.5	0.0211
SI00P	4	6.7	0.1122
UGDH	4	39.3	0.0215
UCHL1	4	41.1	0.0222
SPP1	4	89.3	0.0227
TRIM2	4	154.7	0.0231
FGB	4	156	0.0235
FGG	4	156	0.0441
SDHA	5	0.251	0.0243
PDCD6	5	0.305	0.0245
CCT5	5	10.3	0.0446
PTPRF	5	14.2	0.0248
RPL37	5	40.8	0.0251
ENC1	5	74	0.0336
QP-C	5	132.2	0.0466
SPINK1	5	147.2	0.0256
CANX	5	179.2	0.0263
SOX4	6	21.7	0.0321
HDGF	6	22.6	0.0362
RPS10	6	34.6	0.0177
RPL10A	6	35.4	0.0369
VEGF	6	43.7	0.0372
OSF-2	6	45.4	0.0173
FSCN1	7	5.3	0.0378
CYCS	7	24.9	0.0381
CBX3	7	25.9	0.0289
IGFBP3	7	45.7	0.0389
CLDN4	7	72.7	0.0403
HSPB1	7	75.5	0.0433
CALR	7	92.7	0.0425
COL1A2	7	93.6	0.0457
ATPSJ2	7	98.7	0.0475
AKR1B10	7	133.6	0.0481

Table 1 (continued)

Gene symbol	Chro.	Distance from p arm of each chromosome (Mb)	α
RPS20	8	56.7	0.0482
TCEB1	8	74.6	0.0486
LAPTM4B	8	98.5	0.0497
RPL30	8	98.7	0.0054
KCIP-1	8	101.6	0.0093
PABPC1	8	101.78	0.0119
EEF1D	8	144.4	0.0121
TSTA3	8	144.5	0.0122
RPL8	8	145.6	0.0128
TRA1	9	117.1	0.0136
RPL35	9	121.1	0.0133
HSPA5	9	121.5	0.0135
LCN2	9	124.4	0.0137
DPP7	9	133.4	0.0139
PFKP	10	3.2	0.0223
AKR1C1	10	5.1	0.0146
PLAU	10	75.6	0.0356
DSP	10	76.7	0.0289
TALDO1	11	0.434	0.0143
SLC22A1L	11	2.9	0.0151
TSSC3	11	2.9	0.0611
RPL27A	11	8.7	0.0156
ST5	11	8.8	0.0162
LDHA	11	18.5	0.0168
MDK	11	46.4	0.0162
DOC-1R	11	67.5	0.0167
MMP12	11	102.8	0.0177
HYOU1	11	118.9	0.0183
SCNN1A	12	6.3	0.0185
LDHB	12	21.7	0.0193
KRT7	12	52.3	0.0196
KRT5	12	52.6	0.0197
KRT6E	12	52.6	0.0201
ERBB3	12	56.2	0.0212
NACA	12	56.8	0.0218
TM4SF3	12	71.2	0.0401
NTS	12	86.2	0.0215
ASCL1	12	103.3	0.0219
TXNRD1	12	104.6	0.0223
CKAP4	12	106.6	0.0124
COX6A1	12	120.7	0.0435
BGN	12	122.5	0.0235
RAN	12	129.88	0.0238
RPL36A	14	48.1	0.0243
PGD	14	50.7	0.0248
THBS2	15	37.5	0.0251
TRAF4	15	38.3	0.0253
SPINT1	15	38.7	0.0254
RPL17	15	45.26	0.0411
PKM2	15	70.1	0.0258
IDH2	15	88.2	0.0211
RPL23A	16	0.377	0.0264
MSLN	16	0.753	0.0366
UBE2I	16	1.3	0.0271
RPS2	16	1.95	0.0281
CLDN9	16	3.1	0.0329
ARL6IP	16	18.7	0.0412
EIF3S8	16	28.3	0.0336
TUFM	16	28.9	0.0377
ALDOA	16	30.1	0.038
NME4	16	53.6	0.0381
GPR36	16	57.4	0.0386
CDH1	16	68.5	0.0289
NQO1	16	69.5	0.0396
SLC7A5	16	87.6	0.0397
APRT	16	88.6	0.0411
GALNS	16	88.6	0.0255
RPL13	16	89.3	0.0431
MCP	17	32.4	0.0465

Table 1 (continued)

Gene symbol	Chro.	Distance from p arm of each chromosome (Mb)	α
ERBB2	17	35.11	0.0483
JUP	17	39.8	0.0495
CRF	17	40.39	0.0505
RPL27	17	41.1	0.0046
NME1	17	46.59	0.0082
COL1A1	17	48.6	0.0108
ABCC3	17	49.1	0.0326
NME2	17	49.6	0.0111
RPL38	17	72.7	0.0117
SMT3H2	17	73.6	0.0119
SYNGR2	17	76.6	0.0122
LGALS3BP	17	77.4	0.0127
P4HB	17	80.3	0.0126
PPAP2C	19	0.221	0.0228
GPI	19	39.55	0.0145
HPN	19	40.2	0.0129
ZNF146	19	41.4	0.0131
SPINT2	19	43.4	0.0238
PSMD8	19	43.5	0.0132
YIFIP	19	43.5	0.0135
RPS16	19	44.6	0.0144
CEACAM5	19	46.9	0.0145
CEACAM6	19	46.9	0.0143
GIPR	19	50.8	0.0259
SNRPD2	19	50.9	0.0413
KDELRI	19	53.6	0.0152
RPL28	19	60.6	0.0156
RPS5	19	63.6	0.0267
TRIM28	19	63.7	0.0158
DAP	20	35.6	0.0166
TOP1	20	40.3	0.0172
UBE2C	20	45.1	0.0174
RPS21	20	61.6	0.0268
EEF1A2	20	62.8	0.0185
TFF3	21	42.6	0.0186
TFF1	21	42.7	0.0192
CSTB	21	44.1	0.0201
MIF	22	22.6	0.0202
XBPI	22	27.5	0.0204
PRDX4	X	22.9	0.0198
SYN1	X	46.3	0.0204
TIMPI	X	46.3	0.0209
PLP2	X	47.8	0.0212
MAGED1	X	50.3	0.0331
RPS4X	X	71	0.0124
SSR4	X	152.6	0.0232

corresponding increases in both DNA copy number and mRNA provides further evidence that these could be oncogenes, the activation of which is reflected by genomic amplification and consequent increases in transcript level in lung adenocarcinoma cell lines.

Specific inhibition of EEF1A2 and KCIP-1 expression by siRNAs led to decreased cell proliferation and induction of apoptosis

To further prove the oncogenic function of the identified genes in lung tumorigenesis, we used siRNAs to inhibit the endogenous expression of EEF1A2 and KCIP-1 protein in four lung cancer cell lines (H1563, H229, H522, and SK-LU). Transfection of the cancer cells with specific siRNAs reduced the level of EEF1A2 and KCIP-1 protein by 70–90% 48 h after transfection

(Supplementary Figure 4S). In contrast, EEF1A2 and KCIP-1 protein levels remained unchanged in mock-treated control cells and in cells transfected with a scrambled siRNA sequence. At 48 h after siRNA transfection, the percentage of proliferation of the transfected cancer cells was reduced to 15–30% as compared with 91–100% of cell proliferation of the same cell lines treated with PBS or scrambled siRNA (Supplementary Figure 5S). Apoptosis of siRNA-transfected cells was 27–34%, whereas only 4% of the same cell lines treated with PBS or scrambled siRNA showed apoptosis. These results strongly support an oncogenic role for the identified genes in lung cancer and confirm their potential usefulness as therapeutic targets for the disease.

Amplification and protein expression of KCIP-1 and EEF1A2 in lung tissue

To further validate these findings and to assess the possible clinical significance of the four potential putative oncogenes identified from the cell lines, we first applied fluorescence *in situ* hybridization and immunohistochemical analysis, in parallel, to commercially available human lung tissue microarrays (Ambion, Austin, TX, USA) to evaluate the status of two of these four genes in lung cancer tissue specimens. (Commercially available antibodies to PRDX1 or CALR were not suitable for use in immunohistochemical analysis when this report was written.) Overexpression of KCIP-1 and EEF1A2 protein in the tumors was concordant with amplification of the corresponding genes ($P=0.0003$ for KCIP-1 and $P=0.0011$ for EEF1A2). For example, 16 (35%) of the 46 lung adenocarcinomas in the microarray showed amplification of *KCIP-1*, and strong cytoplasmic staining for KCIP-1 protein was seen in 18 tumors (39%) (Figure 2). We next examined whether overexpression of these genes was associated with increased cell proliferation by analysing Ki-67 expression in contiguous sections of the tissue microarrays. Positive Ki-67 expression was found to correlate with positive expression of both KCIP-1 ($P=0.02$) and EEF1A2 ($P=0.01$). To extend these findings, we then studied 11 tissue microarray blocks comprising normal and tumor tissue specimens from 113 patients with pathologic stage I non-small-cell lung cancer who had undergone curative surgery (Wang *et al.*, 2005). Immunohistochemical analysis showed that EEF1A2 was expressed in 32 cases (28%) and KCIP-1 in 29 cases (26%). Univariate and multivariate Cox proportional hazards models were used to detect possible associations between EEF1A2 and KCIP-1 expression and clinicopathologic variables. Expression of EEF1A2 or KCIP-1 was associated with short overall survival time ($P=0.0012$ for EEF1A2 and $P=0.0026$ for KCIP-1) (Supplementary Figure 6S). Age at diagnosis, histologic type of cancer, degree of tumor differentiation, and smoking history were not associated with survival time.

Although only two genes were validated in the lung tissue microarrays (because available antibodies to the other two genes were not suitable for use in

Table 2 Proteins showing significant overexpression in cancer cell lines relative to those in normal bronchial epithelial cell lines and their correlation coefficients with increased DNA copy number or mRNA values*

Acc. no.	Gene ID	Gene	Mw/pI	Description	r with genomic copy changes ^a	r with mRNA changes ^b
Q06830	5052	PRDX1	48.4/5.4	Peroxisiredoxin 1	0.92364	0.91892
Q05639	1917	EEF1A2	50.5/5.7	Eukaryotic translation elongation factor 1 alpha 2	0.90218	0.89456
P27797	811	CALR	61/5.5	Calreticulin	0.84128	0.86434
P63104	7534	KCIP-1	27/6.5	Tyrosine 3-monooxygenase activation protein, zeta	0.84467	0.85499
P07237	5034	P4HB	54/6.2	Procollagen-proline, 2-oxoglutarate 4-dioxygenase	0.91884	0.76786
Q04695	3872	KRT17	48.0/4.9	Keratin 17	0.00236	0.86892
P09211	2950	GSTP1	23.2/4.7	Glutathione S-transferase pi	0.84218	0.69456
P17936	3486	IGFBP-3	31.6/5.8	Insulin-like growth-factor binding protein 3	0.06412	0.16434
P26641	1937	EEF1G	50/6.4	Eukaryotic translation elongation factor 1 gamma	0.00446	0.85549
P08727	3880	KRT19	44.1/5.2	Keratin 19	-0.04884	0.86786
P04792	3315	HSPB1	22/6.5	Heat shock 27 kDa protein 1	0.00364	0.31892
P00558	5230	PGK1	44.5/4.2	Phosphoglycerate kinase 1	0.50402	0.79456
Q01995	5876	TAGLN	22.5/4.3	Transgelin	-0.34128	-0.26434
P08631	3055	JTK9	59.5/6.8	Hemopoietic cell kinase	-0.01446	0.02549
P09382	3956	LGALS1	16/5.5	Galectin-1, galactoside-binding, soluble, 1	0.026623	0.01123
Q92784	8110	DPF3	25.8/4.8	D4, zinc and double PHD fingers, family 3	0.094884	-0.03214
P54257	9001	HAPI	75.5/6.5	Huntington-associated protein 1	0.12364	-0.08108
P05783	3875	KRT18	48/5.3	Keratin 18	0.010218	0.60544
P05787	3856	KRT8	9.2/4.4	Keratin 8	0.041280	0.84566
P00738	3240	HP	55.2/6.2	Haptoglobin	0.044679	-0.14501
P09769	2268	FGR	59.5/5.2	Gardner-Rasheed feline sarcoma viral oncogene homolog	0.031264	-0.13789
P19838	4790	NFKB1	50.4/6.3	Nuclear factor of kappa light gene enhancer in B-cells 1	0.04467	-0.14501
P29034	6273	S100A2	10.9/4.6	S100 calcium-binding protein A2	0.87964	0.243214
Q13105	7709	ZBTB17	87.9/5.3	Zinc-finger and BTB domain containing 17	-0.17636	0.048108
Q00987	4193	MDM2	75.2/4.8	Transformed 3T3 cell double minute 2	-0.19782	-0.50544
P27816	4134	MAP4	111/5.4	Microtubule-associated protein 4	0.25872	-0.05356
P52732	3832	KIF11	119.2/6.2	Kinesin family member 11	-0.25778	-0.53444
P25205	4172	MCM3	90.9/5.5	Minichromosome maintenance deficient 3	0.25644	0.053666
P08631	3055	HCK	59.5/5.7	Hemopoietic cell kinase	0.65533	0.054501
P09237	4316	MMP7	22.6/5.8	Matrix metalloproteinase 7	0.234987	0.876820
P30305	994	CDC25B	64.9/4.5	Cell division cycle 25B	0.045116	0.283214
P50290	998	CDC42	21.3/6.1	Cell division cycle 42 (GTP-binding protein, 25 kDa)	-0.47636	0.088108
P61586	387	RHOA	19.8/6.9	Ras homolog gene family, member A	-0.49782	-0.00544
P63000	5879	RAC1	21.5/6.8	Ras-related C3 botulinum toxin substrate 1	-0.05583	-0.03566
P07437	203068	TUBB	49.6/6.5	Tubulin, beta polypeptide	0.255533	0.145010
P24864	898	CCNE1	47.1/4.3	Cyclin E1	-0.65116	0.232149
P04141	1437	CSF2	16.9/6.3	Colony stimulating factor 2 (granulocyte-macrophage)	-0.64636	-1.28108
P28072	5694	PSMB6	25.3/5.2	Proteasome (prosome, macropain) subunit, beta type, 6	-0.69782	-1.30544
P00352	216	ALDH1A1	54.7/4.3	Aldehyde dehydrogenase 1 family, member A1	-0.75872	0.03356
Q03013	2948	GTM4	25.3/5.0	Glutathione S-transferase M4	-0.78533	0.134501
P63241	1984	EIF5A	10/4.4	Eukaryotic translation initiation factor 5A	-0.97893	-1.44321
Q01469	2171	EFABP	18.0/4.2	Fatty acid-binding protein 5	0.25684	-0.36432

*Only the gene showing statistically significant increased protein expression with increases in both genomic copy number and transcript simultaneously will be considered as potential putative oncogene in lung adenocarcinoma cells. ^ar, Spearman correlation coefficients between proteins and genomic or mRNA values are based on all six cancer cell lines; bold indicates $P < 0.05$, if $r > 0.84000$. Mw, molecular weight; pI, isoelectric point.

immunohistochemical analysis), these findings are consistent with those from our cell lines, demonstrating again that genomic amplification and consequent increases in amounts of transcript may be, at least in part, driving the abundance of proteins in these lung tumors. The association between expression of these genes and that of Ki-67, a known indicator of poor prognosis in lung cancer (Martin *et al.*, 2004), suggests that activation of these genes may be an indicator of tumor aggressiveness. These results also suggest that expression of EEF1A2 and KCIP-1 proteins in stage I non-small-cell lung cancer may be useful as a marker for distinguishing patients with relatively poor prognosis from those who might benefit from adjuvant treatment.

Discussion

Our current study illustrates the power of integrated functional genomic analyses for identifying putative oncogenes and for evaluating their potential clinical significance. Among the four identified oncogenes, three genes (*PRDX1*, *CALR*, and *KCIP-1*) have been implicated in lung tumorigenesis. *PRDX1* is an antioxidant protein involved in regulating cell proliferation, differentiation, and apoptosis. Kim *et al.* (2003) found *PRDX1* expression to be elevated in both lung cancer and adjacent normal lung tissue, suggesting that activation of *PRDX1* may enhance proliferation in lung cancer. *CALR* has a major role in Ca^{2+} binding and the

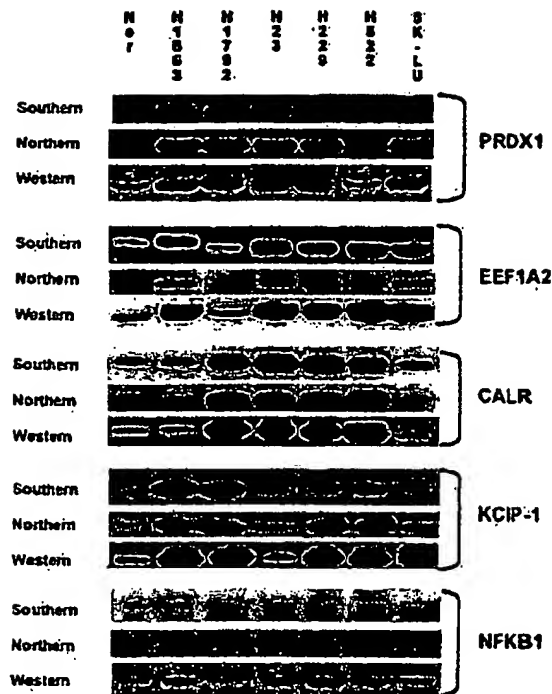


Figure 1 Confirmation by Southern, Northern, and Western blot analyses of increased DNA copies, transcript levels, and protein levels in the four genes identified in high-throughput analyses. For comparison, we arbitrarily chose one gene, *NFKB1*, in which an increased protein level did not correlate with genetic changes. The blotting results are consistent with the results from the CGH array, transcript array, and proteomic analyses. Nor, indicates normal bronchial epithelial cell line. All the experiments were repeated at least three times with each cell line. Means of normalized to β -actin signal intensities on Southern, Northern, and Western blots, along with 95% confidence intervals, were calculated (β -actin signals are not shown in the figure; two different normal bronchial epithelial cell lines were used in the confirmation and only one normal cell line is shown in the figure).

transcriptional regulation of other genes and was recently found to be overexpressed in 73% of 40 lung adenocarcinomas (Oates and Edwards, 2000). *KCIP-1* belongs to the 14-3-3 family, which participates via the MAPK and Wnt signaling pathways in the regulation of many cellular processes including cell proliferation and differentiation as well as tumorigenesis (Thomas et al., 2005). *KCIP-1* was recently found to be expressed in all 12 lung tumors tested in a single-institution study (Qi et al., 2005). Interestingly, *EEF1A2* was originally considered a putative oncogene in ovarian cancer on the basis of its being amplified in 25% and overexpressed in 30% of the same set of ovarian tumors (Anand et al., 2002); functional analyses have established its oncogenic role in cellular transformation (Lee, 2003). Our discovery that *EEF1A2* may be a putative oncogene in lung adenocarcinoma demonstrates the power of our functional genomic strategy for rapidly identifying potential oncogenes.

Although the main focus of this study was to specifically identify putative oncogenes, it should be

noted that 90.7% of the genes showing high protein expression did not show corresponding increases in both DNA copy number and transcript, a finding consistent with that of others that transcriptional, translational, and post-translational regulatory mechanisms can greatly influence the abundance of protein in lung tumorigenesis (Chen et al., 2002). For example, *NFKB1* is a critical arbiter of immune responses, cell survival, and transformation and is often activated in several types of tumors (Chen et al., 2002). Deregulation of *NFKB1* is thought to be modulated through phosphorylation of Ser337 by protein kinase A (Chen et al., 2002). In our study, 68.8% of the genes showing over-representation in the genome did not show elevated transcript levels, implying that at least some of these genes are 'passenger' genes that are concurrently amplified because of their location with respect to amplicons but lack biological relevance in terms of the development of lung adenocarcinoma.

Although the potential oncogenes we identified here are likely to be important, certainly other oncogenes could be involved in the development of lung adenocarcinoma. The oligo microarray we used consists of 22 000 probes, which represent only about 60% of the human genome. Moreover, each probe was designed for the 3' region of expressed sequence tags of the selected genes. Also, our results were initially derived from cancer cell lines, although the findings were later confirmed in human tissue samples. Our ongoing study using microarrays with information on more genes and the development of high-resolution proteomic analyses for use with larger numbers of specimens will allow more comprehensive analyses of the molecular consequences of gene amplifications. Such expanded analyses will very likely lead to the identification of additional oncogenes.

Some of the results of our current study were comparable to those of other studies of lung cancer. For example, genomic copy number and protein levels of *KCIP-1* were previously found to be amplified and overexpressed in primary lung cancers by cDNA clone-based CGH array analysis (Jiang et al., 2004) and proteomic analysis (Chen et al., 2002), respectively. Our functional genomic approach, which integrates simultaneous CGH, transcript microarrays, proteomic analyses, and siRNA, allows us not only to quickly identify potential oncogenes but also to explore their significance as diagnostic and therapeutic targets in tumor progression – more than could be achieved by any technique alone.

Genes identified in this way may serve as promising targets for diagnosis and therapy in lung adenocarcinoma. Further research on the clinical implications of such genes is needed; experiments now underway in our laboratory include overexpression of the genes in normal cells, disruption of the function of these genes in cancer cells, and investigation of how interactions among these genes (or interactions with other known oncogenes) may mediate the expression of the transformed phenotype.

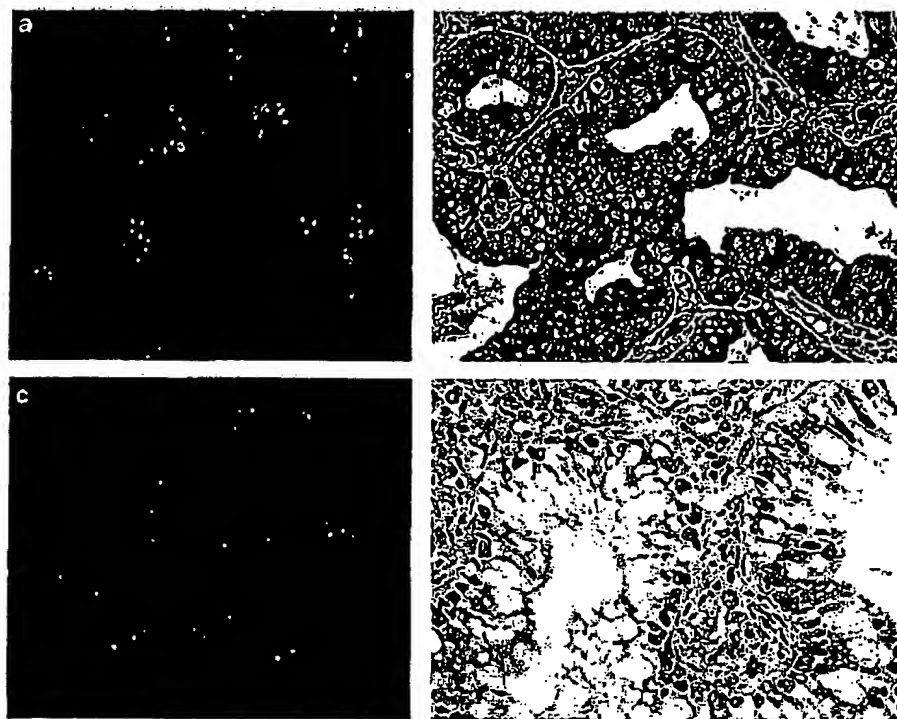


Figure 2 *EEFlA2* amplification is associated with high *EEFlA2* protein expression in lung adenocarcinomas. (a) Cells from a lung adenocarcinoma sample in which *EEFlA2* is amplified show more green signals (*EEFlA2*) than red signals (chromosome 20 centromeric probe) (original magnification, $\times 400$). (b) Immunohistochemical staining of cells from the same tissue sample as in panel a shows strong *EEFlA2* staining in the cytoplasm. (c) A lung adenocarcinoma sample with two copies of *EEFlA2* and chromosome 20 centromeric probe, indicating no *EEFlA2* amplification (original magnification, $\times 400$). (d) Immunohistochemical staining of cells from the same tissue sample as in panel c shows negative staining for *EEFlA2*.

Materials and methods

Cell lines

Six human lung adenocarcinoma cell lines (H23, H229, H1792, SK-LU-1, H522, and H1563) were obtained from the American Type Culture Collection (Manassas, VA, USA). Two normal bronchial epithelial cell lines were obtained from Clontech (Palo Alto, CA, USA). Genomic DNA, mRNA, and protein were derived from a single harvest of these cells.

DNA and RNA profiles by microarray analysis

Genomic DNA labeling and hybridization were performed as described previously (Barrett *et al.*, 2004) with Agilent's Human 1A Oligo Microarray (V2) (Agilent Technologies, Palo Alto, CA, USA), which contains 22 000 unique 60-mer oligos. Details of the protocol for analysing transcripts are available at <http://www.chem.agilent.com>. Map positions for arrayed genes were assigned by identifying the DNA sequence represented in the UniGene cluster and matching it with the Golden Path genome assembly (<http://genome.ucsc.edu/>; Mat 7, 2004 Freeze). Microarray images of DNA copy number and expression were analysed by using AgilentCGH Analytics and Feature Extraction software. DNA copy number profiles that deviated significantly from background signal ratios (measured from normal control cell hybridization, as described elsewhere; Barrett *et al.*, 2004) were interpreted as evidence of true differences in DNA copy number. The criteria for defining genomic over-representation and amplicons are described elsewhere (Hyman *et al.*, 2002); details are given in the

Supplementary Information. An increase in mRNA level was defined as a twofold increase in signal ratio relative to that of the control ($\log_2 > 1$).

Quantitative two-dimensional PAGE and mass spectrometry

Analysis of proteins by two-dimensional PAGE and their identification by mass spectrometry were performed as previously described (Shen *et al.*, 2004). Briefly, protein pellets were solubilized in rehydration buffer, after which the first-dimension isoelectric focusing was carried out with a Protean IEF Cell (Bio-Rad Laboratories) and the second-dimension separation was carried out with Bio-Rad's Ready Gel Precast Gels and the Bio-Rad Criterion Cell apparatus. Protein spots were visualized by silver-based staining, and all gels were assessed with Bio-Rad's PDQuest 2D gel image analysis software. Selected spots were subjected to in-gel tryptic digestion and analysed on a Voyager-DE PRO matrix-assisted laser desorption ionization/time-of-flight mass spectrometer (Applied Biosystems, Foster City, CA, USA). The mass list of the 20 most intense monoisotopic peaks for each sample was entered in the MS-Fit search program (v3.2.1) (<http://prospector.ucsf.edu/ucsfhtml4.0/msfit.htm>) and searched in the National Center for Biotechnology Information protein database.

Southern, Northern, and Western blot analyses

Southern, Northern, and Western blot hybridizations were performed according to standard protocols. cDNA clones for the tested genes were purchased from Invitrogen (Carlsbad,

CA, USA) and prepared as probes for the blot hybridizations. Antibodies used were obtained as follows: PRDX1, CALR, NFKB1, KCIP-1, and β -actin from Santa Cruz Biotechnology (Santa Cruz, CA, USA); and EEF1A2 from Upstate Biotechnology (Waltham, MA, USA).

Fluorescence in situ hybridization and immunohistochemical analyses of lung tissue microarrays

Fluorescence in situ hybridizations and immunohistochemical analyses of KCIP-1 and EEF1A2 were carried out as described elsewhere (Jiang *et al.*, 2002; Wang *et al.*, 2005) with Lung Tissue Microarrays (Ambion, Austin, TX, USA) and 11 homemade microarray blocks containing tissue samples from 113 patients with pathologic stage I non-small-cell lung cancer (Wang *et al.*, 2005). DNA probes specific for KCIP-1 and EEF1A2 were obtained by screening a Human BAC Clone library (Invitrogen) by polymerase chain reaction as described previously (Jiang *et al.*, 2002). The antibodies used for the immunohistochemical analyses were the same as those used for the Western blotting. Cell proliferation of the lung tissues was assessed with a Ki-67 monoclonal antibody from Santa Cruz Biotechnology. Definitions of the cutoff value for a positive result of each antibody are shown in Supplementary Information.

siRNA transfection, cellular proliferation assay, and apoptosis analysis

Transfections were carried out by using siPORT Lipid Transfection Agent (Ambion) with siRNAs targeting KCIP-1 or EEF1A2 or with a scrambled siRNA duplex (siControl) (Dharmacon Inc., Lafayette, CO, USA), with PBS used as a negative control (Jiang *et al.*, 2002). Cells were fixed 24, 48, or 96 h later and subjected to further tests. All siRNAs were prepared by using a transcription-based method with Silencer siRNA according to the manufacturer's instructions (Ambion). Sequences of the individual siRNAs are listed in Supplementary Table 4S. Inhibition of cell growth by the

siRNAs was determined by MTT staining, and cell growth rate was plotted against the percentage of viable cells in the saline-treated controls (a value arbitrarily set at 100%) (Jiang *et al.*, 2002). Apoptosis was analysed by fluorescence cell cycle analysis of terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling with FITC-labeled dUTP (Boehringer Mannheim Biochemicals, Mannheim, Germany) (Jiang *et al.*, 2005).

Statistical analyses

Relationships between gene copy number and mRNA level were examined as described elsewhere (Hyman *et al.*, 2002, Supplementary Information). Correlations between protein abundance and DNA copy number and mRNA expression of the corresponding genes were evaluated with the Spearman correlation coefficient. Fisher's exact test and χ^2 -tests were used to analyse associations between amplification and expression of the candidate genes with various histopathologic variables of the samples in the tissue microarrays. Univariate and multivariate analyses were carried out with Cox's proportional hazards model to determine which independent factors might have a joint significant influence on survival. A *P*-value ≤ 0.05 was considered statistically significant; all statistical tests were based on a two-sided significance level.

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Supplementary Information accompanies the paper on Oncogene website (<http://www.nature.com/onc>)

Variable expression of the translocated *c-abl* oncogene in Philadelphia-chromosome-positive B-lymphoid cell lines from chronic myelogenous leukemia patients

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ABSTRACT The consistent cytogenetic translocation of chronic myelogenous leukemia (the Philadelphia chromosome, Ph¹) has been observed in cells of multiple hematopoietic lineages. This translocation creates a chimeric gene composed of breakpoint-cluster-region (*bcr*) sequences from chromosome 22 fused to a portion of the *abl* oncogene on chromosome 9. The resulting gene product (P210^{c-abl}) resembles the transforming protein of the Abelson murine leukemia virus in its structure and tyrosine kinase activity. P210^{c-abl} is expressed in Ph¹-positive cell lines of myeloid lineage and in clinical specimens with myeloid predominance. We show here that Epstein-Barr virus-transformed B-lymphocyte lines that retain Ph¹ can express P210^{c-abl}. The level of expression in these B-cell lines is generally lower and more variable than that observed for myeloid lines. Protein expression is not related to amplification of the *abl* gene but to variation in the level of *bcr-abl* mRNA produced from a single Ph¹ template.

Chronic myelogenous leukemia (CML) is a disease of the pluripotent stem cell (1). In greater than 95% of patients, the leukemic cells contain the cytogenetic marker known as the Philadelphia chromosome, or Ph¹ (2). This reciprocal translocation event between the long arms of chromosomes 9 and 22 has been used as a disease-specific marker for diagnosis and evaluation of therapy. Multiple hematopoietic lineages, including myeloid and B-lymphoid, contain Ph¹ in early or chronic phase, as well as in the more acute accelerated and blast crisis phases of the disease.

One molecular consequence of Ph¹ is the translocation of the chromosomal arm containing the *c-abl* gene on chromosome 9 into the middle of the breakpoint-cluster region (*bcr*) gene on chromosome 22 (3-6). Although the precise translocation breakpoints are variable, an RNA-splicing mechanism generates a very similar 8-kilobase (kb) mRNA in each case (5-9). The hybrid *bcr-abl* message encodes a structurally altered form of the *abl* oncogene product, called P210^{c-abl} (10-13), with an amino-terminal segment derived from a portion of the exons of *bcr* on chromosome 22 and a carboxyl-terminal segment derived from a major portion of the exons of the *c-abl* gene on chromosome 9. The chimeric structure of *bcr-abl* and the resulting P210^{c-abl} is similar to the structure of the Abelson murine leukemia virus *gag-abl* genome and resulting P160^{v-abl} transforming gene product. Both proteins have very similar tyrosine kinase activities (10, 11, 14) which can be distinguished by their relative stability to denaturing detergents and by their ATP requirements from the recently described tyrosine kinase activity of the *c-abl* gene product (15).

In concert with structural modification of the amino-terminal portion of the *abl* gene, increased level of expression has been implicated in activation of *c-abl* oncogenic potential. Myeloid and erythroid cell lines and clinical samples derived from acute-phase CML patients contain about 10-fold higher levels of the 8-kb *bcr-abl* mRNA and P210^{c-abl} than the *c-abl* mRNA forms (6 and 7 kb) and P145^{c-abl} gene product (5, 8, 9, 11). The higher level of expression of the chimeric *bcr-abl* message in acute-phase cells is not likely to be solely due to the presence of the *bcr* promoter sequences at the 5' end of the gene, since the normal 4.5-kb and 6.7-kb *bcr*-encoded mRNA species are expressed at an even lower level than the normal *c-abl* messages (5, 6).

We have analyzed a series of Epstein-Barr virus-immortalized B-lymphoid cell lines derived from CML patients (16). With such *in vitro* clonal cell lines, we can evaluate whether the presence of Ph¹ always results in synthesis of the chimeric *bcr-abl* message and protein, and whether the quantitative expression varies for cells of B-lymphoid lineage as compared to previously examined myeloid cell lines. Our results show that cell lines that retain Ph¹ do express *bcr-abl* message and protein, but that the level is generally lower and more variable than previously seen for myeloid cell lines. The demonstration that the Ph¹ chromosomal template can vary in its level of expression of P210^{c-abl} suggests that secondary mechanisms, beyond the translocation itself, contribute to the regulation of the *bcr-abl* gene in different cell types or subclones that derive from the affected stem cell.

MATERIALS AND METHODS

Cells and Cell Labelings. Epstein-Barr virus-transformed B-lymphoid cell lines were established from peripheral blood samples of chronic- and acute-phase CML patients as reported (16). The cell lines are designated according to patient number, karyotype, and lineage. For example, SK-CML7Bt(9,22)-33 refers to CML patient 7, B-lymphoid cell line, 9;22 translocation (Ph¹), cell line 33; and SK-CML7BN-2 refers to B-cell line 2 with a normal karyotype derived from the same patient. Repeat karyotype analysis was performed to verify the retention of Ph¹ just prior to analysis for *abl* protein and RNA. Cells were maintained in RPMI 1640 medium with 20% fetal bovine serum. We have not observed any consistent pattern of *in vitro* growth rate that correlates to the stage of disease at the time of transformation with Epstein-Barr virus. Cells (1.5×10^7) were washed twice with Dulbecco's modified Eagle's medium lacking phosphate and

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Abbreviations: *bcr*, breakpoint-cluster region; CML, chronic myelogenous leukemia; kb, kilobase(s).

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populations that lack the Ph¹ marker, supports a secondary or complementary role for Ph¹ in the progression of the disease (24, 25). This chromosome marker is found in chronic, accelerated, and blast-crisis phases of the disease. It is likely that Ph¹ confers some growth advantage, since cells with the marker chromosome eventually predominate the marrow and peripheral blood even in chronic phase. During the phase of blast crisis, many patients develop additional chromosome abnormalities, including duplication of Ph¹, a variety of trisomies, and complex translocations (26). This is suggestive evidence for Ph¹ being a necessary but not sufficient genetic change for the full evolution of the disease.

The realization that one molecular result of Ph¹ is the generation of a chimeric *bcr-abl* protein with functional characteristics and structure analogous to the *gag-abl* transforming protein of the Abelson murine leukemia virus strengthens the argument for an important role of Ph¹ in the pathogenesis of CML. Although the Abelson virus is generally considered a rapidly transforming retrovirus, its effects can range from overcoming growth factor requirements, to cellular lethality, to induction of highly oncogenic tumors in a number of hematopoietic cell lineages (27, 28). Even in the transformation of murine cell targets, there are several lines of evidence that suggest that the growth-promoting activity of the *v-abl* gene product is complemented by further cellular changes in the production of the malignant-cell phenotype (29–31).

The regulation of *bcr-abl* gene expression is complex because the 5' end of the gene is derived from the non-*abl* sequences, *bcr*, normally found on chromosome 22 (6). The level of stable message for the normal *bcr* gene and the normal *abl* gene are both much lower than the level of the *bcr-abl* message and protein from cell lines and clinical specimens derived from myeloid blast-crisis patients (5, 6, 11). Therefore, the high level of *bcr-abl* expression cannot simply be attributed to the regulatory sequences associated with *bcr*. Possibly, creation of the chimeric gene disrupts the normal regulatory sequences and results in a higher level of expression. Variation in *bcr-abl* expression may result from secondary changes in the structure of the chimeric gene or function of *trans*-acting factors that occur during evolution of the disease. Our analysis of P210^{c-abl} and the 8-kb mRNA in Epstein-Barr virus-transformed Ph¹-positive B-cell lines demonstrates that stable message and protein levels from the *bcr-abl* gene can vary over a wide range. This variation does not result from a change in the number of *bcr-abl* templates secondary to gene amplification but more likely from changes in either transcription rate or mRNA stability. We suspect this range of *bcr-abl* expression is not limited to lymphoid cells. Analysis of peripheral blood leukocytes derived from an unusual CML patient who has been in chronic phase with myeloid predominance for 16 years showed a level of P210^{c-abl} one-fifth that of P145^{c-abl}, as detected by metabolic labeling with [³²P]orthophosphate and immunoprecipitation (S.C., O.N.W., and P. Greenberg, unpublished observations). Lower levels of expression of the chimeric mRNA have been demonstrated in clinical samples from chronic-phase CML patients compared to acute-phase CML patients (9). Others have reported chronic-phase patients with variable but, in some cases, relatively high levels of the *bcr-abl* mRNA (32). The sampling variation and the heterogeneous mixture of cell types in clinical samples complicate such analyses. Further work is needed to evaluate whether there is a defined change in P210^{c-abl} expression during the progression of CML. It is interesting to note that among the limited sample of Ph¹-positive B-cell lines we have examined (Table 1), we have seen higher levels of P210^{c-abl} in those derived from patients at more advanced stages of the disease.

It will be important to search for cell-type-specific mechanisms that might regulate expression of *bcr-abl* from Ph¹.

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Genetic Instability in Epithelial Tissues at Risk for Cancer

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ABSTRACT: Epithelial tumors develop through a multistep process driven by genomic instability frequently associated with etiologic agents such as prolonged tobacco smoke exposure or human papilloma virus (HPV) infection. The purpose of the studies reported here was to examine the nature of genomic instability in epithelial tissues at cancer risk in order to identify tissue genetic biomarkers that might be used to assess an individual's cancer risk and response to chemopreventive intervention. As part of several chemoprevention trials, biopsies were obtained from risk tissues (i.e., bronchial biopsies from chronic smokers, oral or laryngeal biopsies from individuals with premalignancy) and examined for chromosome instability using *in situ* hybridization. Nearly all biopsy specimens show evidence for chromosome instability throughout the exposed tissue. Increased chromosome instability was observed with histologic progression in the normal to tumor transition of head and neck squamous cell carcinomas. Chromosome instability was also seen in premalignant head and neck lesions, and high levels were associated with subsequent tumor development. In bronchial biopsies of current smokers, the level of ongoing chromosome instability correlated with smoking intensity (e.g., packs/day), whereas the chromosome index (average number of chromosome copies per cell) correlated with cumulative tobacco exposure (i.e., pack-years). Spatial chromosome analyses of the epithelium demonstrated multifocal clonal outgrowths. In former smokers, random chromosome instability was reduced; however, clonal populations appeared to persist for many years, perhaps accounting for continued lung cancer risk following smoking cessation.

KEYWORDS: chromosome instability; epithelial cells; aerodigestive tract; chemoprevention; cancer risk

THE NEED FOR BIOMARKERS OF CANCER RISK AND RESPONSE TO INTERVENTION

Epithelial cancers remain a major health challenge in the world. Despite improvements in staging and the application and integration of surgery, radiotherapy, and chemotherapy, the 5-year survival rate for individuals with lung cancer is only about 15%.¹ Even if strategies for early detection are successful and lung cancers are detected at a stage where local tumor resection and treatment is curative, these patients will still be at significant risk for developing second primary tumors

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associated with the problem of field cancerization.² Similarly, for individuals with a first head and neck primary tumor, even if the first malignancy is successfully treated, the risk of developing a second primary in the tobacco smoke-exposed field is approximately 40%.³ Similar cancer risk estimates exist for individuals who exhibit severe dysplasia in premalignant epithelial lesions.⁴ For these reasons, it is important to focus on chemopreventive strategies to prevent the development of epithelial malignancies.

Several problems confront chemoprevention trials designed to identify efficacious agents.⁵ First, chemoprevention trials with cancer incidence as a primary endpoint require tens of thousands of subjects and tens of years of intervention and follow-up for statistical evaluation. For example, a recently reported trial involved 30,000 subjects and required 10 years in order to examine the impact of prevention strategies on lung cancer development, only to find a possible increased lung cancer incidence in current smokers who received β -carotene.⁶

The problem of large, long-term trials results from the difficulty in identifying individuals at highest cancer risk who might best benefit from chemopreventive intervention. For example, 20 pack-year smokers, while known to be at relatively increased risk for developing lung cancer, have approximately a 10% lifetime risk for developing lung cancer.⁷ This seriously limits the number of potentially useful strategies that can be clinically explored. A second problem facing chemoprevention trials is that little is known about what agents are likely to have efficacy, and even less is known regarding proper doses, schedules, and durations of treatment. Part of the reason for this problem is that too little is known about the physiologic processes that drive epithelial cancer development.

In order to reduce the number of subjects and the time required to carry out chemoprevention trials and thus allow the exploration of multiple prevention strategies, two types of advances are necessary. First, it is important to identify individuals at significantly increased cancer risk who might best benefit from different types of intervention. Second, in order to allow the rapid identification of agents, doses, and schedules of potentially efficacious agents, it is necessary to identify and validate surrogate endpoints of response that indicate whether the agents are having a positive impact on the target tissue during the chemopreventive intervention.

One approach to identifying individuals at increased aerodigestive tract cancer risk is to explore epidemiologic features of potential subjects. Molecular epidemiologic studies are beginning to identify intrinsic host factors that place some individuals at increased cancer risk, especially those with a chronic smoking history.⁸ Most intrinsic factors identified thus far reflect levels of carcinogen metabolism, repair capabilities of the host following DNA damage, and other measures of intrinsic cellular sensitivity to mutagens. While these factors can provide statistically significant risk ratios in case-control studies that are controlled for tobacco exposure, the detected risk ratios usually fall in the range of 1.5 to 10. Unfortunately, this is not sufficient for the individualization of treatment and is not sufficiently high to significantly reduce the numbers of subjects required for chemoprevention trials with cancer incidence as the primary endpoint.

Another approach to identifying individuals at increased cancer risk is to directly examine the target tissue of individuals with known carcinogen exposure (e.g., chronic tobacco smoke exposure), who have evidence of target organ dysfunction

(e.g., chronic obstructive pulmonary disease, changes in voice quality), or who have clinical evidence of premalignancy (e.g., bronchial metaplasia/dysplasia, oral leukoplakia/erythroplakia, cervical intraepithelial neoplasia). The conventional standard for assessing cancer risk in these situations is the degree of histological change. However, while individuals who show moderate to severe dysplasia are known to be at increased cancer risk when compared to individuals with lesser histologic changes, it is often difficult to distinguish reactive changes to carcinogenic insult from initiated and progressing lesions. Similarly, upon cessation of carcinogenic insult, histologic changes may reverse yet cancer risk may continue for many years. For example, while smoking cessation is associated with decreased bronchial metaplasia,⁹ increased lung cancer risk continues for many years beyond smoking cessation.¹⁰ In fact, nearly half the newly diagnosed lung cancer cases in the USA occur in former smokers.¹¹

The development of assays to identify individuals at high epithelial cancer risk and to directly assess response to intervention in the target tissue is therefore an important research goal. Such assays should be objective and easily quantifiable and, if possible, minimally invasive. Moreover, they should reflect both the disease process and the targeted pathway and thereby be useful in assessing risk and monitoring response to intervention as well as directly testing the hypothesized mechanism of action of the chemopreventive strategy.

In the chemoprevention setting it is important to recognize that one does not know the location of the future cancer. Thus, assays must necessarily be carried out on random biopsies of the field at risk. Even if there are clinically evident premalignant lesions, this does not mean that this is the likely site for a future malignancy. For example, nearly half of the cancers that develop in individuals with oral leukoplakia arise away from the original index lesion. Similarly, since many newly diagnosed lung cancers arise in the peripheral parts of the lung (e.g., adenocarcinomas), especially in former smokers, and since endobronchoscopy predominantly accesses central components of the lung, it is important to identify biomarkers that can reflect global processes ongoing in the target epithelial field associated with increased cancer risk. Their discovery requires a better understanding of the tumorigenesis process in epithelial fields at cancer risk.

THE RATIONALE FOR STUDYING GENOMIC INSTABILITY AS A MARKER OF RISK

Tumors of the aerodigestive tract have been proposed to reflect a "field cancerization" process whereby the whole tissue is exposed to carcinogenic insult (e.g., tobacco smoke) and is at increased risk for multistep tumor development.^{12,13} Several types of clinical and laboratory data support this notion, including the frequent occurrence of synchronous primary and subsequent second primary tumors in the aerodigestive tract (frequently exhibiting dissimilar histologies as well as distinct genetic signatures¹⁴⁻¹⁶) and the presence of premalignant lesions that precede and/or accompany the tumor in the exposed tissue field.¹⁷ The notion of a multistep tumorigenesis process is further supported by serial clinical and histologic evaluations of

target tissue or exfoliated cells where increasing degrees of histological abnormalities are observed over time.¹⁸

A working model for aerodigestive tract tumorigenesis is illustrated in FIGURE 1. Tumorigenesis in the face of carcinogenic exposure likely involves a chronic process of tissue injury and wound healing. DNA damage induced by the carcinogen is likely fixed into permanent genetic changes (e.g., chromosome damage, chromosome non-disjunction, gene mutation, gene deletion, etc.) during the process of proliferation. This damage would be expected to be distributed throughout the exposed tissue field leading to a background of generalized genomic damage (depicted in FIGURE 1 as a background mat of increasing density). Chronic injury and repair likely leads to the accumulation of cells with increasing amounts of genetic changes as well as the outgrowth of abnormal clones (triangles in FIGURE 1) carrying an accumulation of genetic changes important for selective survival, dysregulated growth, and preferential epithelial take-over by initiated clones (see FIGURE 2).

Cellular and molecular evidence for the field carcinogenesis and multistep tumorigenesis model comes from many laboratories.^{19,20} With the advent of a wide array of molecular technologies, a large number of specific molecular genetic and epigenetic changes involving specific oncogenes, tumor suppressor genes, cell regulatory genes, and repair genes have now been described for aerodigestive tract cancers. The identification of these specific molecular changes have now provided probes to explore specific events occurring in premalignant lesions adjacent to aerodigestive tract tumors.²¹⁻²⁴ Frequently, these premalignant lesions showed a subset of the same molecular changes found in the associated tumor, suggesting that these lesions might represent precursor lesions for the associated tumors (i.e., a manifestation of

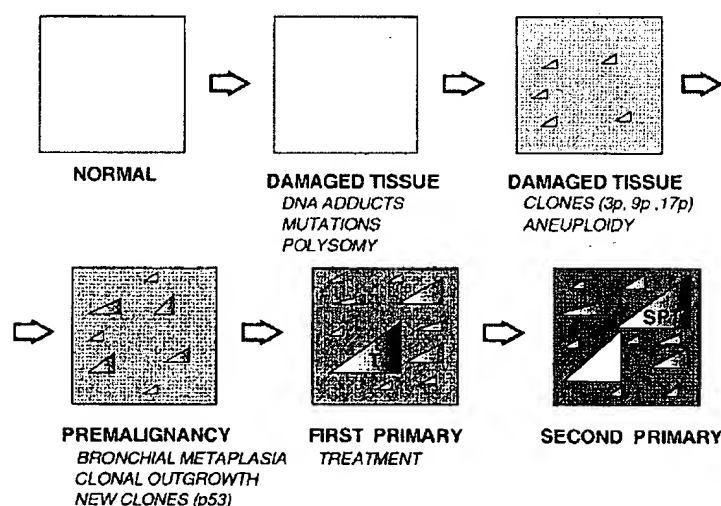


FIGURE 1. Field cancerization and multistep tumorigenesis.

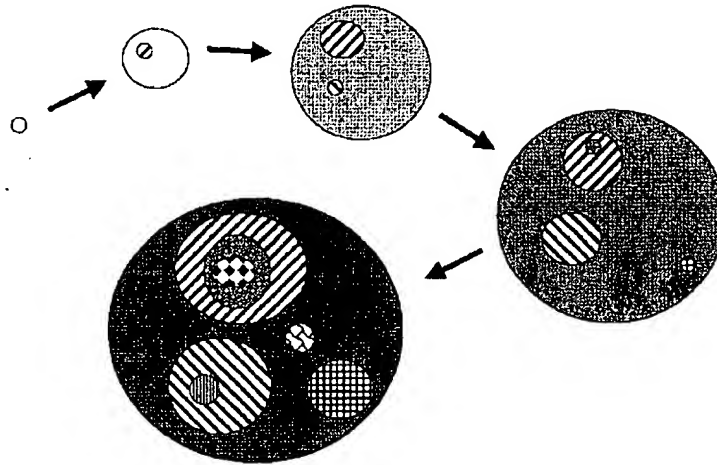


FIGURE 2. Multiple focal clonal evolution during multistep tumorigenesis.

a multistep tumorigenesis process). For example, studies of the premalignant lesions adjacent to head and neck tumors have provided evidence for a gradual accumulation of genetic alterations accompanied by evidence for dysregulation of cellular control mechanisms (e.g., alterations in expression of PCNA, EGFR, TGF- β , p53, and cyclin D1).²⁵⁻²⁸

These types of studies have now also been applied to the target epithelium of individuals at increased risk for aerodigestive tract cancer (i.e., individuals with a chronic smoking/alcohol history and/or prior aerodigestive tract cancer). Several groups (using polymerase chain reaction, PCR, analysis of microdissected epithelium) have now demonstrated the presence of clonal outgrowths in the target premalignant epithelium of individuals at increased risk for cancer.²⁹⁻³¹ For example, examination of bronchial biopsies derived from individuals with a 20 pack-year smoking history demonstrated that 76% of the cases showed evidence for LOH (3p14, 9p21, or 17p13) in at least one of six lung biopsy sites. On a per site basis, some form of LOH was observed in 25% of the sites examined.²⁹

If aerodigestive tract cancer development reflects a field cancerization process involving multistep events, then risk and response information should be able to be derived from random biopsies or exfoliated cells from the field at risk or from assessments of tissue undergoing similar processes. Hypothetically, lesions exhibiting the greatest degree of genomic instability, clonal outgrowth, and abnormal epithelial regulation would be at the highest relative aerodigestive tract cancer risk. Similarly, an active chemopreventive intervention might be expected to decrease these manifestations of risk. Reduced risk manifestations include decreased levels of ongoing genetic instability, decreased frequency of clonal outgrowths, and increased epithelial growth regulation.

THE MEASUREMENT OF CHROMOSOME INSTABILITY USING CHROMOSOME *IN SITU* HYBRIDIZATION

Molecular genetic techniques, while extremely useful for detecting clonal changes in target tissues, are somewhat limited in their ability to detect random genetic instability. Conventional cytogenetic assays are useful for detecting chromosome instability and clonal chromosome changes. However, they require numbers of dividing cells for karyotypic analysis that are difficult to attain in the setting of biopsies acquired during the course of a chemoprevention trial. A technique was therefore needed that would allow chromosome instability measurements in situations where few cells are available (e.g. small biopsies, brushings, or sputum samples) and where the target material might be fixed. It was also desirable to have a technique that would be adaptable to tissue sections, whereby spatial information could be retained and genotype/phenotype associations could be determined on the same or adjacent tissue sections. The technique of *in situ* hybridization (ISH) involves the use of DNA probes that recognize either chromosome-specific repetitive target sequences, chromosome single gene copy sequences, or sequences along the whole chromosome length or chromosome segments.³² We have adapted the ISH technique for formalin-fixed, paraffin-embedded tissue sections and have applied it to a variety of tissues, including the aerodigestive tract.^{33,34}

Using probes that label the centromere regions of specific chromosomes, this assay permits determination of the average chromosome number per cell for each specimen. This assay is also useful for detecting generalized chromosome instability during the tumorigenesis process. Normal diploid populations should have two copies of each autosomal chromosome and should rarely show three or more chromosome copies per cell (chromosome polysomy), especially in tissue sections where nuclear truncation results in an under-representation of chromosome copy number. Thus, the detection of cells with three or more chromosome copies would indicate the presence of chromosome instability.

To examine this technique's potential for characterizing the multistep tumorigenesis process in the aerodigestive tract, we measured the fraction of cells exhibiting three or more chromosome copies in apparently contiguous epithelial transitions from normal to hyperplastic to dysplastic to carcinomas, all on a single tissue slice of head and neck squamous cell carcinomas.³⁴ In these specimens, greater than 35% of the cases of adjacent "normal" epithelium, greater than 65% of the cases of hyperplastic epithelium, and greater than 95% of the dysplastic and tumor regions showed evidence of chromosome polysomy. Of interest, similar transitions of chromosome instability were observed with at least four different chromosome probes. Similar trends have also been observed in amenable tissue from other epithelial malignancies, including cervix, bladder, and breast.³⁵ These results thus suggested that the notions of field cancerization and multistep tumorigenesis might apply to several epithelial tissues and that measures of chromosome instability might be useful for monitoring this process.

In the situations described above, the premalignant lesions examined might be considered to represent epithelium at 100% risk of being in a cancer field, since they were located in the adjacent epithelium to the cancer. This then raises the question of the nature of genetic instability in the epithelium of individuals at increased risk

for developing cancer. To explore this issue, we obtained biopsies during the course of leukoplakia chemoprevention trials exploring the use of 13-*cis*-retinoic acid in reversing leukoplakia and probed them for genetic instability using *in situ* hybridization. In one retrospective study and in one prospective study of subjects with oral leukoplakia, the results indicate that those subjects whose pretreatment biopsies harbor relatively high levels of genomic instability (i.e., more than 3% of the cells examined showing at least 3 chromosome 9 copies per cell) have a significantly higher likelihood of suffering early onset of head and neck cancer.^{36,37} Interestingly, half of the tumors that did develop occurred away from the biopsy site used to measure genetic instability. This result suggests that genomic instability measurements in carcinogen-exposed tissue can provide useful cancer risk estimates.

THE RELATIONSHIP BETWEEN TOBACCO EXPOSURE AND CHROMOSOME INSTABILITY

In recent years, the aerodigestive tract chemoprevention group at M.D. Anderson Cancer Center has initiated three sequential biomarker-associated chemoprevention trials involving chronic smokers with a greater than 20 pack-year smoking history. In each of these studies, endobronchial biopsies were obtained from six defined sites within the lung, including the carina and at bifurcation points at the upper, middle, and lower right lung and at the upper and lower left lung. Biopsies were obtained prior to and following chemopreventive intervention and were subjected to *in situ* hybridization analysis in addition to analyses for other biomarkers. The first important finding was that some degree of chromosome polysomy was evident in all lung sites examined, and this was observed independently of the particular chromosome probe utilized.³⁸ This finding supports the notion that random chromosome changes may be occurring throughout the exposed lung field.

In a second study, bronchial biopsies were obtained from individuals with a 20 pack-year smoking history. In this study, most of the subjects involved were current smokers.³⁹ Interestingly, all cases who showed metaplasia at one of six biopsy sites also showed chromosome polysomy in at least one biopsy site; overall, 88% of the sites showed some evidence of chromosome 9 polysomy.⁴⁰ Evidence for genetic instability was also detected in patients who did not show evidence of bronchial metaplasia in any of six biopsy sites despite a strong smoking history. In fact, more than 90% of the cases and more than 60% of the sites showed significant chromosome polysomy (i.e., at least three copies in at least 2 % of the cells examined). These results suggest that the lungs of long-term smokers show significant evidence of genetic instability, and this instability can be detected throughout the accessible bronchial tree, even when bronchial metaplasia is not evident.

These studies in current smokers has allowed us to examine the relationship between the levels of genetic instability detected and subject characteristics such as smoking status (current or former), smoking history, and lung tissue pathologic changes. Evaluable biopsy material has now been obtained from more than 108 current smokers, including more than 480 evaluable biopsy sites. The mean metaplasia index in these current smokers was 30.4%. For the total population studied, the median chromosome index for the bronchial biopsies was 1.41 (range, 1.04–1.61)

and the median chromosome polysomy index was 2.0% (range 0–8.7%). This can be compared to a mean chromosome index between 1.2–1.4 for lymphocytes and very rare chromosome polysomy. Interestingly, the intrasubject variability in chromosome instability was relatively low in most subjects and was less than the intersubject variability. These results suggested that chronic smokers harbor detectable chromosome instability throughout the accessible bronchial tree (supporting the field carcinogenesis notion) and that information from one biopsy site might yield representative information for the rest of the lung field.

Since most of the current smokers exhibited bronchial metaplasia in at least one of the biopsied sites, this allowed us to examine the relationship between chromosome instability and histologic changes, both on a site-by-site basis and on a per case basis. On a site-by-site basis, the chromosome indices of lesions showing squamous metaplasia were similar to those not showing metaplasia (i.e., median 1.43 vs. 1.43), and the degree of chromosome polysomy in metaplastic lesions were only slightly higher than in non-metaplastic sites (medians: 2.2% vs. 1.8%, respectively). Thus, the presence or absence of squamous metaplasia at a biopsy site does not necessarily correlate with the degree of underlying genomic instability. On the other hand, those subjects with metaplasia indices of at least 15% also showed higher levels of chromosome polysomy than did subjects with metaplasia index below 15% (medians: 2.4% vs. 1.8%, $p = 0.005$). Thus, these chromosome instability assessments in current smokers appeared to reflect a more global process in the lung field.

Tobacco exposure has been shown to significantly increase the risk of developing lung cancer, and the degree of risk is related to the extent of tobacco exposure. We were interested in determining the relationship between individuals' smoking history parameters and the levels of chromosome change found in their lungs following years of tobacco exposure. While there was significant intersubject variation for similar tobacco exposure histories, overall there was a significant correlation between the degree of chromosome polysomy and the intensity of ongoing tobacco exposure (packs/day, $p = 0.02$ on a per site basis) and with the extent of tobacco exposure (pack-years, $p = 0.003$). Thus the amount of chromosome polysomy reflects the intensity and extent of tobacco exposure. At the same time, individuals with similar smoking histories showed widely divergent amounts of chromosome polysomy, possibly reflecting differences in intrinsic sensitivity between subjects. There was also strong correlation between the chromosome index and the duration of the smoking history (smoking years) and total accumulated exposure (pack-years, $p = 0.0001$). These results suggest that tobacco exposure is associated with the initiation and accumulation of chromosome instability in the exposed lung; however individuals are differentially sensitive to carcinogenic insult. The working hypothesis is that those individuals who accumulate the highest degree of chromosome changes will be at the highest lung cancer risk.

Many of the bronchial biopsies from chronic smokers examined by *in situ* hybridization showed a rise in the chromosome index above that expected for a diploid cell population, especially in subjects with an extensive smoking history. The rise in chromosome index was also accompanied by an increase in the fraction of cells exhibiting at least 3 chromosome copies per cell. To determine if a rise in the tissue chromosome index was due to clonal expansion of populations with chromosome trisomy, the chromosome copy number and relative coordinates of each cell scored in

the bronchial epithelium was recorded and a spatial genetic map was created.⁴¹ We then developed algorithms for calculating localized chromosome indices within the tissue. Since trisomic clones would have, on average, three chromosomes instead of two, those cells involved in neighborhoods with chromosome indices three-halves that of diploid populations could be marked as being part of a trisomic clone. Similarly, groups of cells with chromosome indices half that of diploid populations could be marked as being part of a monosomic clone. This allowed the generation of a second-order, two-dimensional genetic map representation of the bronchial epithelium showing the relative locations of cells involved in monosomic and trisomic clonal outgrowths. When adjacent tissue sections from the same bronchial biopsy were probed separately for different chromosomes, the detected clones appeared to occupy separate subregions of the epithelium. This result suggests that not only are the lungs of chronic smokers undergoing a process of genetic instability, they are experiencing the outgrowth of multiple clones throughout the exposed lung field, as postulated by the models shown in FIGURES 1 and 2. One advantage of this clonal approach is that the contribution of both monosomic and multisomic clones can be detected.

Since smoking cessation has been suggested to reduce the lung cancer risk, it was of interest to determine whether the levels of chromosome instability would decrease following smoking cessation. This question was possible to examine because our third sequential chemoprevention trial involved subjects who had discontinued smoking. So far, more than 220 subjects (more than 650 biopsies) who have quit smoking (mean 9.9 quit-years) have been evaluated for chromosome instability in their lungs. Despite the fact that the mean metaplasia index in this group is 5.8% (considerably less than that in current smokers), chromosome instability is still observed in the majority of subjects.⁴² While the mean chromosome polysomy level is reduced to 1.0%, some individuals continue to show polysomy levels above 5%. Interestingly, while the overall chromosome polysomy levels were reduced in these individuals who stopped smoking, the mean chromosome index remained at about 1.4 with some individuals exhibiting chromosome indices as high as 1.8. Initial chromosome mapping studies suggest that while random chromosome instability seems to decrease following smoking cessation, the clonal outgrowths may remain for many years in the lung. The working hypothesis is that those individuals who show the greatest degree of remaining chromosome instability are at the highest lung cancer risk despite smoking cessation. Long-term follow-up on these subjects will be necessary to test this hypothesis.

SUMMARY AND CONCLUSIONS

Aerodigestive tract tumorigenesis appears to be a multistep process taking place throughout the tissue fields of exposure. When viewed in the context of chromosome changes, carcinogen exposure appears to be associated with the random acquisition of chromosome polysomy throughout the exposed field, the degree of which is related to the degree and extent of carcinogen exposure as well as to the intrinsic susceptibility of the exposed individual. Continued exposure leads to continued acquisition of new changes and, in association with chronic wound-healing processes, to the

accumulation of clonal outgrowths throughout the target tissue. Although the ultimate malignancy may occur in only one or few tissue sites, manifestations of the instability process that drives tumorigenesis is globally present in the tissue. Thus random biopsies may provide useful risk information for the exposed field as a whole. Even when carcinogen exposure is reduced or chemopreventive strategies are initiated and histologic manifestations of the tumorigenesis process subside, the genetic scars of prior exposure remain in the form of clonal outgrowths and may explain continued lung cancer risk in ex-smokers. Future chemoprevention strategies need to focus on reducing the degree of chromosome instability and on trying to eliminate residual abnormal clonal outgrowths in the aerodigestive tract. In this setting, the measurement of chromosome instability in the target tissue will be useful in assessing cancer risk as well as response to intervention.

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- ☐ **GRAY SCALE DOCUMENTS**
- ☒ **LINES OR MARKS ON ORIGINAL DOCUMENT**
- ☐ **REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY**
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